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NEWS	1			Web Page for STN Seminar Schedule - N. America
NEWS NEWS		AUG	06 06	CAS REGISTRY enhanced with new experimental property tags FSTA enhanced with new thesaurus edition
NEWS		AUG		CA/CAplus enhanced with additional kind codes for granted
MENO		100	13	patents
NEWS		AUG		CA/CAplus enhanced with CAS indexing in pre-1907 records
NEWS	6 1	AUG	27	Full-text patent databases enhanced with predefined
			0.77	patent family display formats from INPADOCDB
NEWS NEWS		AUG		USPATOLD now available on STN CAS REGISTRY enhanced with additional experimental
MEMO	0 1	100	20	spectral property data
NEWS	9 5	SEP	07	STN AnaVist, Version 2.0, now available with Derwent
				World Patents Index
NEWS			13	FORIS renamed to SOFIS
NEWS		SEP		INPADOCDB enhanced with monthly SDI frequency
NEWS	12 3	SEP	17	CA/CAplus enhanced with printed CA page images from 1967-1998
NEWS	13 3	SEP	17	CAplus coverage extended to include traditional medicine
				patents
NEWS		SEP		EMBASE, EMBAL, and LEMBASE reloaded with enhancements
NEWS	15 (OCT	02	CA/CAplus enhanced with pre-1907 records from Chemisches Zentralblatt
NEWS	16 (OCT	19	BEILSTEIN updated with new compounds
NEWS			15	Derwent Indian patent publication number format enhanced
NEWS			19	WPIX enhanced with XML display format
NEWS			30	ICSD reloaded with enhancements
NEWS			04	LINPADOCDB now available on STN
NEWS NEWS			14 17	BEILSTEIN pricing structure to change USPATOLD added to additional database clusters
NEWS			17	IMSDRUGCONF removed from database clusters and STN
NEWS			17	DGENE now includes more than 10 million sequences
NEWS		DEC		TOXCENTER enhanced with 2008 MeSH vocabulary in
				MEDLINE segment
NEWS			17	MEDLINE and LMEDLINE updated with 2008 MeSH vocabulary
NEWS		DEC		CA/CAplus enhanced with new custom IPC display formats
NEWS	28 1	DEC	17	STN Viewer enhanced with full-text patent content from USPATOLD
NEWS	EXPRI	ESS	CUI	SEPTEMBER 2007: CURRENT WINDOWS VERSION IS V8.2, RRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP), D CURRENT DISCOVER FILE IS DATED 19 SEPTEMBER 2007.
NEWS	HOURS	S		N Operating Hours Plus Help Desk Availability
	LOGII	И		Lcome Banner and News Items
NEWS	IPC8		For	general information regarding STN implementation of IPC 8

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 COST IN U.S. DOLLARS
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 ENTRY
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 FULL ESTIMATED COST
 0.21
 0.21

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STRUCTURE FILE UPDATES: 26 DEC 2007 HIGHEST RN 959588-76-2 DICTIONARY FILE UPDATES: 26 DEC 2007 HIGHEST RN 959588-76-2

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TSCA INFORMATION NOW CURRENT THROUGH June 29, 2007

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http://www.cas.org/support/stngen/stndoc/properties.html

Uploading C:\Program Files\Stnexp\Queries\10560862.str

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chain nodes:
7 8 10 11 12 16
ring nodes:
1 2 3 4 5 6
chain bonds:
2-8 3-7 5-10 6-16 10-11 11-12
ring bonds:
1-2 1-6 2-3 3-4 4-5 5-6
exact/norm bonds:
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normalized bonds:
1-2 1-6 2-3 3-4 4-5 5-6
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G1:C,S

G2:C,O

Match level: 1:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 10:CLASS 11:CLASS 12:CLASS 16:CLASS 16:CLASS 6eneric attributes: 7:

Saturation : Unsaturated 8: Saturation : Unsaturated

L1 STRUCTURE UPLOADED

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Structure attributes must be viewed using STN Express query preparation.

4 ANSWERS

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=> s 11 SAMPLE SEARCH INITIATED 09:47:44 FILE 'REGISTRY' SAMPLE SCREEN SEARCH COMPLETED -6907 TO ITERATE

29.0% PROCESSED 2000 ITERATIONS INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED)

FULL FILE PROJECTIONS: ONLINE **COMPLETE** **COMPLETE** BATCH

PROJECTED ITERATIONS: 133158 TO 143122 PROJECTED ANSWERS: 53 TO

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SEARCH TIME: 00.00.01

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=> s 12
L3 4 L2
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=> d 1-4 ibib abs

L3 ANSWER 1 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:962046 CAPLUS

DOCUMENT NUMBER: 143:266952

TITLE: Preparation of bipyridyl amides as modulators of

metabotropic glutamate receptor-5
INVENTOR(S): Bonnefous, Celine; Kamenecka, Theodore M.; Vernier,

Jean-Michel

PATENT ASSIGNEE(S): Merck & Co., Inc., USA SOURCE: PCT Int. Appl., 79 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

									APPLICATION NO.								
					A1 20050901			WO 2005-US3952									
	W:	AE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,
		NO.	NZ,	OM,	PG.	PH.	PL,	PT.	RO,	RU,	SC,	SD,	SE,	SG,	SK.	SL,	SY.
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AU	2005						2005	0901		AU 2	005-	2153	79		2	0050	209
CA	2555	402			A1		2005	0901		CA 2	005-	2555	402		2	0050	209
EP	1715	867			A1		2006	1102		EP 2	005-	7131	11		2	0050	209
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CN	1933	838			A		2007	0321		CN 2	005-	8000	4732		2	0050	209
JP	2007	5246	82		Т		2007	0830		JP 2	006 -	5531	89		2	0050	209
IN	2006	DN04	346		A		2007	0713		IN 2	006-	DN43	46		2	0060	727
US	2007	1495	47		A1		2007	0628		US 2	006-	5894	07		2	0060	811
PRIORIT											004-						
										WO 2	005-	US39	52		W 2	0050	209
OTHER S	OURCE	(S):			CAS	REAC	T 14	3:26	6952	; MA	RPAT	143	:266	952			

AB The title compds. I [X = N, C; Y = N, C, C(halo); R1 = H, alkyl, cycloalkyl, etc.; R2 = H, alkyl, aryl, etc.; R3 = aryl, halo, alkyl, etc.; R2 and R3 may be joined together with the atoms to which they are attached to form a (un)saturated 4-7 membered ring containing 0-2 heteroatoms selected

from

O, S and N; R4 = aryl, heteroaryl, halo, etc.] which are mGluR5 modulators useful in the treatment or prevention of diseases and conditions in which mGluR6 is involved, including but not limited to psychiatric and mood disorders such as schizophrenia, anxiety, depression, bipolar disorders, and panic, as well as in the treatment of pain, Parkinson's disease, cognitive dysfunction, epilepsy, circadian rhythm and sleep disorders, such as shift-work induced sleep disorder and jet-lag, drug addiction, drug abuse, drug withdrawal, obesity and other diseases, were prepared Thus, amidation of pyridin-2-amine with 3-amino-5,6-diphenylpyrazine-2-carboxylic acid afforded the amide II. The exemplified compds. I have mGluR5 inhibitory activity as shown by inhibition at 10 μM or less in the calcium flux assay or 100 μM or less or less in the PI assay. The invention is also directed to pharmaceutical compns. comprising compds. I.

invention is also directed to pharmaceutical compns. comprising compds. I.

REFERENCE COUNT: 5 THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 2 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2004:1127366 CAPLUS

DOCUMENT NUMBER: 142:56362

TITLE: Preparation of 3-substituted 5,6-diaryl-pyrazine-2-

carboxamide and 2-sulfonamide derivatives as

cannabinoid receptor 1 (CB1) modulators

SOURCE: PCT Int. Appl., 120 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PA:	PATENT NO.					KIND DATE			APPLICATION NO.									
WO	WO 2004111034				A1 20041223			WO 2004-SE970					2	0040	616			
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
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MARPAT 142:56362

II

AB Title compds. I [wherein R1, R2 = independently (un) substituted Ph, thienyl, pyridinyl; R3 = X-Y-NR5R6; X = absent, CO, or SO2; Y = absent, NH optionally substituted by an alkyl group; R5, R6 = independently (un)substituted amino/alkyl, (CH2)r(phenyl)s, (un)saturated 5-8-membered heterocyclyl; R5 = H and R6 = defined above; or R5NR6 = (un)substituted (un)saturated 5-8-membered heterocycly1; r = 0-4; s = 1 when r = 0, otherwise s = 1 or 2; R5NR6 = (un)substituted (un)saturated 5-8-membered heterocyclyl; R4 = (CH2) nCO2R7; n = 0-4; R7 = (un) substituted cycloalkyl/cyclo/alkyl, (CH2) nphenyl, saturated or partially unsatd. 5-8-membered heterocyclyl, CONH2 and derivs.; n = defined as above; and pharmaceutically acceptable salts thereofl were prepared as cannabinoid 1 (CB1) receptor modulators. For example, reacting 3-(tert-butoxycarbonyl)-5,6-bis(4-chlorophenyl)pyrazine-2-carboxylic acid (preparation given) with tert-butylhydrazine hydrochloride gave pyrazine II. I are active at the CB1 receptor (IC50 < 1 μ M), most preferred compds. have IC50 < 200 nM. For instance, II exhibited an IC50 (hCB1) = 1.8 nM. Thus, I and their pharmaceutical compns. are useful for the treatment of obesity, psychiatric and neurol. disorders (no data). THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS REFERENCE COUNT: RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L3 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1980:41887 CAPLUS DOCUMENT NUMBER: 92:41887

ORIGINAL REFERENCE NO.: 92:6993a,6996a

TITLE: Chemistry of diaminomaleonitrile. 5. Dihydropyrazine synthesis

AUTHOR(S): Ohtsuka, Yozo; Tohma, Eiko; Kojima, Sigeru; Tomita, Nobuo

CORPORATE SOURCE: Sagami Chem. Res. Cent., Sagamihara, 229, Japan SOURCE: Journal of Organic Chemistry (1979), 44(26), 4871-6 CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal
LANGUAGE: English
OTHER SOURCE(S): CASREACT 92:41887

AB Condensation of RCHO (R = optionally substituted Ph) with Schiff bases I (R1 = optionally substituted Ph, CHMe2) in the presence of NEt3 <20° is accompanied by regiospecific hydration of the nitrile groups to give 3-cyanoacrylamide derivs. II, which cyclize readily into 1,2-dihydropyrazines III and IV. The substituent effect on the product ratio is examined, and the reaction mechanism is discussed in terms of a new general reaction pattern of diaminomaleonitrile derivative Reactions of III and IV by oxidation, reduction, hydantoin formation with isocyanates, and cyanoethylation are also reported.

L3 ANSWER 4 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1956:69468 CAPLUS

DOCUMENT NUMBER: 50:69468

ORIGINAL REFERENCE NO.: 50:13047b-i,13048a-b

TITLE: Pteridines. XIV. Further studies on a new approach to

pteridine synthesis

AUTHOR(S): Taylor, E. C., Jr.; Garland, Robert B.; Howell,

Charles F.

CORPORATE SOURCE: Univ. of Illinois, Urbana

SOURCE: Journal of the American Chemical Society (1956), 78,

210-13

CODEN: JACSAT: ISSN: 0002-7863

DOCUMENT TYPE: Journal

LANGUAGE: Unavailable

OTHER SOURCE(S): CASREACT 50:69468

AB cf. C.A. 50, 2608h. 3-Amino-5,6-diphenylpyrazinamide (I) (1.509 g.) and 10 cc. BzCl refluxed 4 h., cooled, and diluted with 250 cc. petr. ether gave 1.179 g. 2,6,7-triphenyl-4(3H)-pteridinone (II), white needles, m. 290° (from CHZCl2-petr. ether and then aqueous HCONMe2) (all m.ps. are

corrected). The N-PhCH2 derivative (III) of I (0.5 g.) and 25 cc. AcCl

refluxed 4

h. and diluted with 25 cc. petr. ether yielded 0.36 g. 3-acetylamino-5,6-diphenylpyrazinamide (IV), bright yellow platelets, m. 207-8° (from CRC13-petr. ether). III (0.835 g.), 10 cc. Ac20, and 10 cc. MeCN refluxed 4 h. and evaporated to dryness in vacuo, and the residue treated with EtOH and evaporated to dryness again gave 0.472 g. N-PhCH2 derivative (V) of IV, tan crystals, m. 149-50° (from CH2C12-petr. ether). V (0.613 g.)

refluxed 3 h. with 0.5 g. Na in 10 cc. absolute EtOH and poured into 50 cc. H2O gave 0.503 g. III, m. 186-7°. 3-PhCH2 derivative of II gave similarly 93% III. I (2.53 g.), 5 cc. PhNCO, and 25 cc. dry pyridine refluxed 1 h. and cooled yielded 2.81 g. 3-(3-phenylureido)-5,6-diphenylpyrazinamide (VI), light yellow platelets, m. 240,5-1.5°

cc. dry pyridine refluxed 2 h., cooled, treated with C, and diluted with petr. ether gave 1.03 g. N-PhCH2 derivative (VII) of VI, sparkling white platelets, m. 210° (from aqueous EtOH). VI (0.523 g.) and 7 g. polyphosphoric acid (VIII) heated 2 h. at 150° (CO2 was evolved), and diluted with 50 cc. H2O, and the precipitate sublimed at 200° and 2 mm. gave 0.134 g. I, m. 204-5°; the sublimation residue sublimed at 300° and 2 mm, gave 3,5,7-triphenvl-2,4(1H,3H)-pteridinedione (IX), colorless solid, m. 327-8° (decomposition). III and VIII heated 45 min. at 150° gave 52% I and 63% VII. I (0.97 g.), 2 cc. PhNCO, and 10 cc. pyridine refluxed 3 days, cooled, diluted with 40 cc. CH2Cl2 and 250 cc. petr. ether, and filtered, and the filtrate evaporated to dryness gave 0.418 g. IX, white needles, m. 327-8° (decomposition) (from aqueous HCONMe2). III gave similarly 51% IX. I (1.52 g.), 3 cc. PhNCS, and 15 cc. pyridine refluxed 1 h., cooled, and diluted with 150 cc. petr. ether yielded 1.92 g. 3-(3-phenylthioureido) analog (X) of I, light yellow platelets, m. 233° (from aqueous HCONMe2). I (1.67 g.), 3 cc. PhNCS, and 15 cc. pyridine refluxed 3 days, cooled overnight, and filtered gave 1.87 g. 2-mercapto-3,6,7-triphenyl-4(3H)-pteridinone (XI), fine yellow needles, m. 301-2° (sublimed at 250° and 1 mm.). X heated similarly with PhNCS gave also XI. N-Bu derivative of I (2.70 g.), 3.5 cc. PhNCS, and 10 cc. pyridine refluxed 4 days, cooled, and diluted with 20 cc. CH2Cl2 and 100 cc. petr. ether yielded 1.49 g. 2-PhNH analog of XI, pale yellow crystals, m. 323-4° (from aqueous HCONMe2). I (1.34 q.), 2 cc. iso-PrNCS, and 20 cc. pyridine refluxed 2 days, cooled, and diluted with 20 cc. CHCl3 and 100 cc. petr. ether gave 1.05 g. 3-(3-isopropylthioureido) analog (XII) of VI, white platelets, m. 251-2° (from CH2C12-cyclohexane). III (1.04 g.), 1.2 cc. iso-PrNCS, and 15 cc. pyridine refluxed 2 days and poured onto 200 g. ice yielded 0.7 g. N-PhCH2 derivative (XIII) of XII, pale yellow crystals, m. 170° (from 70% AcOH). XII (1.24 q.) refluxed 6 h. with 1 q. Na in 25 cc. absolute EtOH, poured into 100 cc. H2O, and filtered, and the orange solid digested with dilute HCl gave 0.174 g. 2-mercapto-3-isopropyl-6,7-diphenyl-4(3H)pteridinone, light yellow needles, m. 270° (from aqueous EtOH); the filtrate acidified with concentrated HCl gave 0.72 g. 2-isopropylamino-6,7diphenyl-4(3H)-pteridinone (XIV), bright lemon-yellow platelets, m. 324-5° (from aqueous EtOH). XIII (0.390 g.) refluxed 3 h. with 0.1 g. Na in 5 cc. absolute EtOH and poured into 50 cc. H2O yielded 0.30 g. 3-PhCH2 derivative of XIV, sparkling yellow crystals, m. 305-7° (decomposition) (from aqueous HCONMe2). 3-Amino-5,6-diphenylthiopyrazinamide (XV) (1.1 g.) and 10 cc. BzCl refluxed 1.5 h., cooled, diluted with 50 cc. EtOH, refluxed 1 h., and evaporated to dryness, and the residue suspended in hot EtOH and filtered gave 2,6,7-triphenyl-4(3H)-pteridinethione, yellow crystals, m. 323-4° (sublimed). XV (1.23 g.), 3.4 cc. PhNCS, and 10 cc. pyridine refluxed 2 h., cooled, and diluted with 180 cc. petr. ether yielded 2.06 g. compound C47H33N9O (structure tentatively assigned), fine vellow needles, m. 369-70° (from aqueous HCONMe2), also obtained by refluxing the mixture for 3 days. It was recovered in 93% yield after refluxing 43 h. with concentrated HCl. XV (1.04 g.), 2 cc. PhNCS, and 10 cc. pyridine refluxed 36 h., diluted with 150 cc. hot petr. ether, and allowed to stand gave a small amount of unidentified, colorless needles, m. 72-157°, fine yellow needles, and cushions of orange prisms. The fine yellow needles and orange prisms recrystd. from pyridine-petr. ether yielded 1.15 g. 2-anilino-6,7-diphenyl-4(3H)pteridinethione, long yellow needles, m. 261-2°.

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L3 ANSWER 1 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN

IT 863908-32-1P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (USes)

(preparation of bipyridyl amides as modulators of metabotropic glutamate receptor-5)

RN 863908-32-1 CAPLUS

CN Pyrazinecarboxamide, 3-amino-5,6-diphenyl-N-2-pyridinyl- (9CI) (CA INDEX NAME)

L3 ANSWER 2 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN

IT 811441-53-9P, 5,6-Bis(4-chlorophenyl)-N-(3-hydroxypropyl)-N' (piperidin-1-yl)pyrazine-2,3-dicarboxamide
 RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
 (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(Uses)
(drug candidate; preparation of 3-substituted 5,6-diarylpyrazine-2-carboxamide and 2-sulfonamide derivs. as CB1 modulators)

RN 811441-53-9 CAPLUS

CN 2,3-Pyrazinedicarboxamide, 5,6-bis(4-chlorophenyl)-N-(3-hydroxypropyl)-N'-1-piperidinyl- (9CI) (CA INDEX NAME)

- L3 ANSWER 3 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN
- IT 71871-24-4P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 71871-24-4 CAPLUS

CN Pyrazinecarboxamide, 3-cyano-6-(4-cyanophenyl)-5-(4-methylphenyl)- (9CI) (CA INDEX NAME)

- ANSWER 4 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN
- ΙT 859300-58-6P, Pyrazinamide, 3-(3-isopropyl-2-thioureido)-5,6diphenyl-RL: PREP (Preparation)

(preparation of) 859300-58-6 CAPLUS RN

Pyrazinamide, 3-(3-isopropyl-2-thioureido)-5,6-diphenyl- (5CI) (CA INDEX NAME)

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Substance data SEARCH and crossover from CAS REGISTRY in progress... Use DISPLAY HITSTR (or FHITSTR) to directly view retrieved structures.

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2000 ITERATIONS 29.0% PROCESSED INCOMPLETE SEARCH (SYSTEM LIMIT EXCEEDED) SEARCH TIME: 00.00.01

4 ANSWERS

FULL FILE PROJECTIONS: ONLINE **COMPLETE** BATCH **COMPLETE** PROJECTED ITERATIONS: 133158 TO 143122 PROJECTED ANSWERS: 53 TO 499 L5 4 L4

MISSING OPERATOR L5 SSS

The search profile that was entered contains terms or nested terms that are not separated by a logical operator.

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FULL ESTIMATED COST	ENTRY 0.47	SESSION 25.82
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE ENTRY	TOTAL
CA SUBSCRIBER PRICE	0.00	-3.12

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=> file reg COST IN U.S. DOLLARS	SINCE FILE ENTRY	TOTAL
FULL ESTIMATED COST	0.47	26.29
DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)	SINCE FILE ENTRY	TOTAL SESSION
CA SUBSCRIBER PRICE	0.00	-3.12

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STRUCTURE FILE UPDATES: 26 DEC 2007 HIGHEST RN 959588-76-2 DICTIONARY FILE UPDATES: 26 DEC 2007 HIGHEST RN 959588-76-2

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http://www.cas.org/support/stngen/stndoc/properties.html

=> s 11 sss full FULL SEARCH INITIATED 09:51:36 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED = 139072 TO ITERATE

100.0% PROCESSED 139072 ITERATIONS 202 ANSWERS SEARCH TIME: 00.00.01

L6 202 SEA SSS FUL L1

=> file caplus

 COST IN U.S. DOLLARS
 SINCE FILE ENTRY
 TOTAL SESSION 172.10

 FULL ESTIMATED COST
 172.10
 198.39

 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)
 SINCE FILE TOTAL ENTRY SESSION 0.00
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FILE 'CAPLUS' ENTERED AT 09:51:41 ON 27 DEC 2007 USE IS SUBJECT TO THE TERMS OF YOUR STN CUSTOMER AGREEMENT. PLEASE SEE "HELP USAGETERMS" FOR DETAILS. COPYRIGHT (C) 2007 AMERICAN CHEMICAL SOCIETY (ACS)

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FILE COVERS 1907 - 27 Dec 2007 VOL 147 ISS 26 FILE LAST UPDATED: 26 Dec 2007 (20071226/ED)

Effective October 17, 2005, revised CAS Information Use Policies apply. They are available for your review at:

http://www.cas.org/infopolicy.html

=> s 16

L7 42 L6

L7 ANSWER 1 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2007:84319 CAPLUS

DOCUMENT NUMBER: 146:184452

TITLE: Preparation of thioamides as selective CB1 antagonists

for treating obesity, psychiatric and neurol.

INVENTOR(S): Bostrom, Jonas; Cheng, Leifeng; Olsson, Roine PATENT ASSIGNEE(S): Astrazeneca AB, Swed.; Astrazeneca UK Limited

SOURCE: PCT Int. Appl., 44pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

GI

PATENT N				KIN	D	DATE		2	APPL	ICAT	ION	NO.		D.	ATE		
WO 20070				A2		20070125		1	WO 2006-GB2638						20060717		
W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
	CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
	GE,	GH,	GM,	HN,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KM,	KN,	KP,	
	KR,	ΚZ,	LA,	LC,	LK,	LR,	LS,	LT,	LU,	LV,	LY,	MA,	MD,	MG,	MK,	MN,	
	MW,	MX,	MZ,	NA,	NG,	NI,	NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RS,	RU,	
	SC,	SD,	SE,	SG,	SK,	SL,	SM,	SY,	ΤJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	
	US,	UZ,	VC,	VN,	ZA,	ZM,	ZW										
RW:	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	ES,	FI,	FR,	GB,	GR,	HU,	IE,	
	IS,	IT,	LT,	LU,	LV,	MC,	NL,	PL,	PT,	RO,	SE,	SI,	SK,	TR,	BF,	ΒJ,	
	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	SN,	TD,	TG,	BW,	GH,	
	GM,	KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	ΑZ,	BY,	
	KG,	ΚZ,	MD,	RU,	ТJ,	TM											
ITY APPI	LN.	INFO	. :						GB 2	005-	1473	9		A 2	0050	719	

PRIORITY APPLN. INFO.: OTHER SOURCE(S): CASREACT 146:184452; MARPAT 146:184452

$$\begin{bmatrix} R^4 \\ N \\ N \end{bmatrix}_{R} \begin{bmatrix} R^2 \\ N \end{bmatrix}_{R} \begin{bmatrix} R^1 \\ N \end{bmatrix}_{R}$$

AB The title compds. I [HET = II, III, IV, etc. (wherein Rl = alkoxy (optionally substituted by one or more F atoms), O(CH2)pPh, etc.; p = 1-3; m = 0-3; R2 = alkyl, alkoxy, OH, etc.; n = 0-3; R4 = H, alkyl, alkoxy, etc.); R3 = (un)substituted cyclohexyl, piperidino, Ph, etc.], useful in the treatment of obesity, psychiatric and neurol. disorders, were prepared E.g., a multi-step synthesis of 4-(3-(cyclohexylamino)carbonothioyl]-1-(2,4-dichlorophenyl)-4-methyl-1H-pyrazol-5-yl]bhenyl propanel-sulfonate, starting from 4-hydroxyproplophenone, was given. Compds. I are active at the CBI receptor (ICSO < 1 µM). The invention also relates to methods for therapeutic use of compds. I and to pharmaceutical compns. containing them.

IT 921628-24-2P 921628-25-3P 921628-26-4P 921628-27-5P 921628-28-6P 921628-29-7P

921628-30-0P

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of thioamides as CB1 antagonists for treating obesity, psychiatric and neurol. disorders)

RN 921628-24-2 CAPLUS

CN 2-Pyrazinecarbothioamide, 5,6-bis(4-chlorophenyl)-N-(4,4-difluorocyclohexyl)-3-(methoxymethyl)- (CA INDEX NAME)

RN 921628-25-3 CAPLUS

CN 2-Pyrazinecarbothioamide, 5,6-bis(4-chlorophenyl)-N-(2-hydroxycyclohexyl)-3-(methoxymethyl)- (CA INDEX NAME)

RN 921628-26-4 CAPLUS

CN 2-Pyrazinecarbothioamide, 5,6-bis(4-chlorophenyl)-N-[2-(dimethylamino)cyclohexyl]-3-(methoxymethyl)- (CA INDEX NAME)

- RN 921628-27-5 CAPLUS
- CN 2-Pyrazinecarbothioamide, 5,6-bis(4-chlorophenyl)-N-(3-hydroxycyclohexyl)3-(methoxymethyl)- (CA INDEX NAME)

- RN 921628-28-6 CAPLUS
- CN 2-Pyrazinecarbothioamide, N-(3-aminocyclohexyl)-5,6-bis(4-chlorophenyl)-3-(methoxymethyl)- (CA INDEX NAME)

- RN 921628-29-7 CAPLUS
- CN 2-Pyrazinecarbothioamide, 6-(4-chlorophenyl)-N-(4,4-difluorocyclohexyl)-3(methoxymethyl)-5-[4-(3,3,3-trifluoropropoxy)phenyl]- (CA INDEX NAME)

$$\begin{array}{c} \text{C1} \\ \text{F}_3\text{C}-\text{CH}_2-\text{CH}_2-\text{O} \\ \\ \text{N} \\ \text{C}-\text{NH} \end{array}$$

RN 921628-30-0 CAPLUS

CN 2-Pyrazinecarbothioamide, 5,6-bis(4-chlorophenyl)-N-[1-(hydroxymethyl)-3-methylbutyl]-3-(methoxymethyl)- (CA INDEX NAME)

L7 ANSWER 2 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2006:646507 CAPLUS

DOCUMENT NUMBER: 145:271733

AUTHOR(S):

TITLE: Straightforward Access to Pyrazines, Piperazinones, and Quinoxalines by Reactions of 1,2-Diaza-1,3-

butadienes with 1,2-Diamines under Solution,

Solvent-Free, or Solid-Phase Conditions

Aparicio, Domitila; Attanasi, Orazio A.; Filippone, Paolino; Ignacio, Roberto; Lillini, Samuele;

Mantellini, Fabio; Palacios, Francisco; de Santos,

Jesus M.

CORPORATE SOURCE: Istituto di Chimica Organica, Universita degli Studi

di Urbino Carlo Bo, Urbino, 61029, Italy

SOURCE: Journal of Organic Chemistry (2006), 71(16), 5897-5905 CODEN: JOCEAH; ISSN: 0022-3263

PUBLISHER: American Chemical Society

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 145:271733

AB The preparation of tetrahydropyrazines, dihydropyrazines, pyrazines, piperazinones, and quinoxalines by 1,4-addition of 1,2-diamines to 1,2-diaza-1,3-butadienes bearing carboxylate, carboxamide, or phosphorylated groups at the terminal carbon and subsequent internal heterocyclization is described. The solvent-free reaction of carboxylated 1,2-diaza-1,3-butadienes with the same reagents affords piperazinones, while phosphorylated 1,2-diaza-1,3-butadienes yield phosphorylated pyrazines. The solid-phase reaction of polymer-bound 1,2-diaza-1,3-

butadienes with 1,2-diamines produces pyrazines.

907161-24-4P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of pyrazines, piperazinones, and quinoxalines by

1,4-addition/heterocyclization of 1,2-diaza-1,3-butadienes with 1,2-diamines under solution, solvent-free, or solid-phase conditions)

907161-24-4 CAPLUS RN

CN Pyrazinecarboxamide, N,N,3-trimethyl-5,6-diphenyl- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

108 FORMAT

THERE ARE 108 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE

ANSWER 3 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:1103771 CAPLUS

DOCUMENT NUMBER: 143:367331

TITLE: Pyrazine derivatives as adenosine antagonists, their preparation, pharmaceutical compositions, and use in

therapy INVENTOR(S): Tsutsumi, Hideo; Tabuchi, Seiichiro; Minagawa,

Masatoshi; Akahane, Atsushi

Astellas Phama Inc., Japan PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 204 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

		IT NO.				KIND DATE				APPLICATION NO.								
					A1 20051013			WO 2005-JP5663						20050322				
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	, BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
		CN,	co,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	, EC,	EE,	EG,	ES,	FI,	GB,	GD,	
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	, JP,	KE,	KG,	KP,	KR,	KZ,	LC,	
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG.	, MK,	MN,	MW,	MX,	MZ,	NA,	NI,	
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	, sc,	SD,	SE,	SG,	SK,	SL,	SM,	
		SY,	ΤJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	, US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW
	RW:	BW,	GH,	GM,	KΕ,	LS,	MW,	MZ,	NA,	SD,	, SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	
		AZ,	BY,	KG,	KZ,	MD,	RU,	TJ,	TM,	AT,	, BE,	BG,	CH,	CY,	CZ,	DE,	DK,	
		EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	IE,	IS,	, IT,	LT,	LU,	MC,	NL,	PL,	PT,	
		RO,	SE,	SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	, CI,	CM,	GA,	GN,	GQ,	GW,	ML,	
					TD,													
CA	2562	126			A1		2005	1013		CA 2	2005~	2562	126		2	0050	322	
EP	1737	841			A1		2007	0103		EP 2	2005-	7215	90		2	0050	322	
	R:	ΑT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	EE,	, ES,	FI,	FR,	GB,	GR,	HU,	IE,	
		IS,	IT,	LI,	LT,	LU,	MC,	NL,	PL,	PT,	, RO,	SE,	SI,	SK,	TR			
							2007	0328		CN 2	2005-	8001	0591		2	0050	322	
	2007						2007	1101		JP 2	2006-	5294	02		2	0050	322	
IN	2006	CN03					2007	0615		IN 2	2006-0	CN36	09		2	0060	928	
MX	2006	PA11:	247		A		2006	1129		MX :	2006-1	PA11	247		2	0060	929	

KR 2007008674 A 20070117 KR 2006-722911 20061031 PRIORITY APPLN. INFO.: AU 2004-901772 A 20040401 W0 2005-<u>D</u>F5663 W 20050322

OTHER SOURCE(S): MARPAT 143:367331

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* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The invention relates to pyrazine derivs. of formula I, which are adenosine antagonists. In compds. I, R is H or (un) substituted lower alkyl; X is H, halo, OH, SH, cyano, acyl, (un) substituted lower alkyl, (un) substituted lower alkoxy, (un) substituted aryl, etc.; Y is H, halo, OH, SH, cyano, acyl, (un) substituted lower alkyl, (un) substituted lower alkoxy, (un) substituted lower alkylthio, (un) substituted amino, (un) substituted aryl, or (un) substituted heteroaryl; and Z is (un) substituted aryl or (un) substituted heteroaryl; or a salt thereof. The invention also relates to the preparation of I, pharmaceutical compns. containing I, or a pharmaceutically acceptable salt thereof, in admixt. with a pharmaceutically acceptable carrier, as well as to the use of the compns. in the treatment of disorders responding to adenosine antagonists. Oxidation of 2-isopropyl-6-(phenylethynyl)-3-pyridazinone (II) to the corresponding dione followed by condensation with 2,3-diamine-2-butenedinitrile resulted in the formation of pyridazinylpyrazine III, which underwent regioselective substitution with 4-methoxybenzylamine, debenzylation, and hydrolysis to give pyrazinecarboxamide IV. The amide of IV was cleaved followed by decarboxylation, bromination with N-bromosuccinimide, and palladium-catalyzed coupling with 5-ethynyl-1-methyl-1H-imidazole to give pyrazinylpyridazinone V. The tested compds. express high affinity for adenosine receptors, with compound V expressing Ki values of 0.72 nM and 0.25 nM for adenosine A1 and A2a receptors, resp.

G66263-05-0P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3pyridazinyl)-5-phenyl-2-pyrazinecarboxamide 866263-20-9P,
3-Cyano-5-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridazinyl)-6-phenyl-2pyrazinecarboxamide 866263-29-8P, 3-Amino-6-(1-isopropyl-6-oxo1,6-dihydro-3-pyridazinyl)-5-phenyl-2-pyrazinecarbothioamide
866264-95-1P, 3-Amino-5-(2-bromophenyl)-6-(1-isopropyl-6-oxo-1,6dihydro-3-pyridazinyl)-2-pyrazinecarboxamide
RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic
preparation); THU (Therapeutic use); BIOL (Biological study); PREP
(Preparation); RACT (Reactant or reagent); USES (USes)

(drug candidate; preparation of pyrazine derivs. as adenosine antagonists) 866263-05-0 CAPLUS

CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pvridazinyl]-5-phenyl- (9CI) (CA INDEX NAME)

RN

RN

CN Pyrazinecarboxamide, 3-cyano-5-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-6-phenyl- (9CI) (CA INDEX NAME)

- RN 866263-29-8 CAPLUS
- CN Pyrazinecarbothioamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-5-phenyl- (9CI) (CA INDEX NAME)

- RN 866264-95-1 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(2-bromophenyl)-6-[1,6-dihydro-1-(1methylethyl)-6-oxo-3-pyridazinyl]- (9CI) (CA INDEX NAME)

IT 866263-11-8P, 3-Amino-6-(1-ethyl-6-oxo-1,6-dihydro-3-pyridazinyl)5-phenyl-2-pyrazinecarboxamide 866263-15-2P,
3-Amino-6-(1-methyl-6-oxo-1,6-dihydro-3-pyridazinyl)-5-phenyl-2pyrazinecarboxamide 866263-21-0P, 3-Cyano-6-(1-isopropyl-6-oxo1,6-dihydro-3-pyridazinyl)-5-phenyl-2-pyrazinecarboxamide
866263-33-4P, 3-Amino-5-(1-isopropyl-6-oxo-1,6-dihydro-3pyridazinyl)-6-phenyl-2-pyrazinecarboxamide 866263-45-8P,
3-Amino-5-(4-fluorophenyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridazinyl)2-pyrazinecarboxamide 866263-47-0P, 3-Amino-5-(3-fluorophenyl)-6(1-isopropyl-6-oxo-1,6-dihydro-3-pyridazinyl)-2-pyrazinecarboxamide
866263-45-0P, 3-Bis (4-wethoxybenzyl) amino]-6-(1-isopropyl-6-oxo-

1,6-dihydro-3-pyridazinyl)-5-phenyl-2-pyrazinecarboxamide
866264-96-2P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3pyridazinyl)-5-(2-thienyl)-2-pyrazinecarboxamide 866264-97-3P,
3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridazinyl)-5-(4-pyridyl)-2pyrazinecarboxamide 866264-98-4P, 3-Amino-6-(1-isopropyl-6-oxo1,6-dihydro-3-pyridazinyl)-5-(6-methoxy-3-pyridyl)-2-pyrazinecarboxamide
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)

(drug candidate; preparation of pyrazine derivs. as adenosine antagonists) 866263-11-8 CAPLUS

CN Pyrazinecarboxamide, 3-amino-6-(1-ethyl-1,6-dihydro-6-oxo-3-pyridazinyl)-5phenyl- (9CI) (CA INDEX NAME)

RN

RN 866263-15-2 CAPLUS

CN Pyrazinecarboxamide, 3-amino-6-(1,6-dihydro-1-methyl-6-oxo-3-pyridazinyl)-5-phenyl- (9CI) (CA INDEX NAME)

RN 866263-21-0 CAPLUS

CN Pyrazinecarboxamide, 3-cyano-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pvridazinyl]-5-phenyl- (9CI) (CA INDEX NAME)

RN

CN Pyrazinecarboxamide, 3-amino-5-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridazinyl]-6-phenyl- (9CI) (CA INDEX NAME)

- RN 866263-45-8 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-5-(4-fluorophenyl)- (9CI) (CA INDEX NAME)

- RN 866263-47-0 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-5-(3-fluorophenyl)- (9CI) (CA INDEX NAME)

- RN 866263-55-0 CAPLUS
- CN Pyrazinecarboxamide, 3-[bis[(4-methoxyphenyl)methyl]amino]-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-5-phenyl- (9CI) (CA INDEX NAME)

- RN 866264-96-2 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-5-(2-thienyl)- (9CI) (CA INDEX NAME)

- RN 866264-97-3 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-5-(4-pyridinyl)- (9CI) (CA INDEX NAME)

- RN 866264-98-4 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-5-(6-methoxy-3-pyridinyl)- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

1 ANSWER 4 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:1078246 CAPLUS

DOCUMENT NUMBER: 143:367330

TITLE: Pyrazine derivatives as adenosine antagonists, their preparation, pharmaceutical compositions, and use in

therapy

INVENTOR(S): Tsutsumi, Hideo; Tabuchi, Seiichiro; Minagawa, Masatoshi; Akahane, Atsushi

PATENT ASSIGNEE(S): Fujisawa Pharmaceutical Co. Ltd., Japan

SOURCE: U.S. Pat. Appl. Publ., 54 pp.

CODEN: USXXCO DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE		
US 2005222159	A1	20051006	US 2005-87761	20050324		
US 7265120	B2	20070904				
PRIORITY APPLN. INFO.:			AU 2004-901772 A	20040401		
OTHER SOURCE(S):	MARPAT	143:367330				

GI

* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT *

The invention relates to pyrazine derivs. of formula I, which are AB adenosine antagonists. In compds. I, R is H or (un) substituted lower alkyl; X is H, halo, OH, SH, cyano, acyl, (un) substituted lower alkyl, (un) substituted lower alkoxy, (un) substituted aryl, etc.; Y is H, halo, OH, SH, cyano, acyl, (un) substituted lower alkyl, (un) substituted lower alkoxy, (un) substituted lower alkylthio, (un) substituted amino, (un) substituted aryl, or (un) substituted heteroaryl; and Z is (un) substituted aryl or (un) substituted heteroaryl; or a salt thereof. The invention also relates to the preparation of I, pharmaceutical compns. containing I, or a pharmaceutically acceptable salt thereof, in admixt. with a pharmaceutically acceptable carrier, as well as to the use of the compns. in the treatment of disorders responding to adenosine antagonists. Oxidation of 2-isopropyl-6-(phenylethynyl)-3-pyridazinone (II) to the corresponding

dione followed by condensation with 2,3-diamine-2-butenedinitrile resulted in the formation of pyridazinylpyrazine III, which underwent regioselective substitution with 4-methoxybenzylamine, debenzylation, and hydrolysis to give pyrazinecarboxamide IV. The amide of IV was cleaved followed by decarboxylation, bromination with N-bromosuccinimide, and palladium-catalyzed coupling with 5-ethynyl-1-methyl-1H-imidazole to give pyrazinylpyridazinone V. The tested compds. express high affinity for adenosine receptors, with compound V expressing Ki values of 0.72 nM and

0.25 nM for adenosine A1 and A2a receptors, resp. 866263-05-0P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3pyridazinyl)-5-phenyl-2-pyrazinecarboxamide 866263-20-9P, 3-Cyano-5-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridazinyl)-6-phenyl-2pyrazinecarboxamide 866263-29-8P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridazinyl)-5-phenyl-2-pyrazinecarbothioamide 866264-95-1P, 3-Amino-5-(2-bromophenyl)-6-(1-isopropyl-6-oxo-1,6dihydro-3-pyridazinyl)-2-pyrazinecarboxamide RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); RACT (Reactant or reagent); USES (Uses) (drug candidate; preparation of pyrazine derivs. as adenosine antagonists)

866263-05-0 CAPLUS RN CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-

pyridazinyll-5-phenyl- (9CI) (CA INDEX NAME)

RN 866263-20-9 CAPLUS CN

Pyrazinecarboxamide, 3-cyano-5-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridazinyl]-6-phenyl- (9CI) (CA INDEX NAME)

RN 866263-29-8 CAPLUS

Pyrazinecarbothioamide, 3-amino-6-[1.6-dihydro-1-(1-methylethyl)-6-oxo-3-CN pyridazinyl]-5-phenyl- (9CI) (CA INDEX NAME)

- RN 866264-95-1 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(2-bromophenyl)-6-[1,6-dihydro-1-(1methylethyl)-6-oxo-3-pyridazinyl]- (9CI) (CA INDEX NAME)

866263-11-8P, 3-Amino-6-(1-ethyl-6-oxo-1,6-dihydro-3-pyridazinyl)-5-phenv1-2-pyrazinecarboxamide 866263-15-2P, 3-Amino-6-(1-methyl-6-oxo-1,6-dihydro-3-pyridazinyl)-5-phenyl-2pyrazinecarboxamide 866263-21-0P, 3-Cyano-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridazinyl)-5-phenyl-2-pyrazinecarboxamide 866263-33-4P, 3-Amino-5-(1-isopropyl-6-oxo-1,6-dihydro-3pyridazinyl)-6-phenyl-2-pyrazinecarboxamide 866263-45-8P, 3-Amino-5-(4-fluorophenyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridazinyl)-2-pvrazinecarboxamide 866263-47-0P 866263-55-0P, 3-[Bis(4-methoxybenzyl)amino]-6-(1-isopropyl-6-oxo-1,6-dihydro-3pyridazinyl)-5-phenyl-2-pyrazinecarboxamide 866264-96-2P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridazinyl)-5-(2-thienyl)-2pyrazinecarboxamide 866264-97-3P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridazinyl)-5-(4-pyridyl)-2-pyrazinecarboxamide 866264-98-4P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3pvridazinvl)-5-(6-methoxv-3-pvridvl)-2-pvrazinecarboxamide RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (drug candidate; preparation of pyrazine derivs. as adenosine antagonists) 866263-11-8 CAPLUS

RN 866263-11-8 CAPLUS
CN Pyrazinecarboxamide, 3-amino-6-(1-ethyl-1,6-dihydro-6-oxo-3-pyridazinyl)-5phenyl- (9C1) (CA INDEX NAME)

- RN 866263-15-2 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-(1,6-dihydro-1-methyl-6-oxo-3-pyridazinyl)-5-phenyl- (9CI) (CA INDEX NAME)

- RN 866263-21-0 CAPLUS
- CN Pyrazinecarboxamide, 3-cyano-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-5-phenyl- (9CI) (CA INDEX NAME)

- RN 866263-33-4 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-6-phenyl- (9CI) (CA INDEX NAME)

- RN 866263-45-8 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-5-(4-fluorophenyl)- (9CI) (CA INDEX NAME)

- RN 866263-47-0 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-5-(3-fluorophenyl)- (9CI) (CA INDEX NAME)

- RN 866263-55-0 CAPLUS
- CN Pyrazinecarboxamide, 3-[bis[(4-methoxyphenyl)methyl]amino]-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-5-phenyl- (9CI) (CA INDEX NAME)

- RN 866264-96-2 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-5-(2-thienyl)- (9CI) (CA INDEX NAME)

- RN 866264-97-3 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-5-(4-pyridinyl)- (9CI) (CA INDEX NAME)

- RN 866264-98-4 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridazinyl]-5-(6-methoxy-3-pyridinyl)- (9CI) (CA INDEX NAME)

L7 ANSWER 5 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2005:962046 CAPLUS

DOCUMENT NUMBER: 143:266952

TITLE: Preparation of bipyridyl amides as modulators of

metabotropic glutamate receptor-5

INVENTOR(S): Bonnefous, Celine: Kamenecka, Theodore M.: Vernier, Jean-Michel

PATENT ASSIGNEE (S): Merck & Co., Inc., USA PCT Int. Appl., 79 pp. SOURCE:

CODEN: PIXXD2 DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

APPLICATION NO. PATENT NO. KIND DATE DATE WO 2005079802 A1 20050901 WO 2005-US3952 20050209 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GO, GW, ML, MR, NE, SN, TD, TG AU 2005215379 20050901 AU 2005-215379 A1 20050209 CA 2555402 20050901 CA 2005-2555402 20050209 A1 EP 1715867 20061102 EP 2005-713111 A1 20050209 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, FI, RO, CY, TR, BG, CZ, EE, HU, PL, SK, IS CN 1933838 20070321 CN 2005-80004732 Α 20050209 JP 2006-553189 JP 2007524682 Τ 20070830 20050209 IN 2006DN04346 IN 2006-DN4346 Α 20070713 US 2007149547 A1 20070628 US 2006-589407 PRIORITY APPLN. INFO.: US 2004-544627P P 20040212

CASREACT 143:266952; MARPAT 143:266952

WO 2005-US3952

W 20050209

OTHER SOURCE(S):

- AB The title compds. I [X = N, C; Y = N, C, C(halo); R1 = H, alkyl, cycloalkyl, etc.; R2 = H, alkyl, aryl, etc.; R3 = aryl, halo, alkyl, etc.; R2 and R3 may be joined together with the atoms to which they are attached to form a (un)saturated 4-7 membered ring containing 0-2 heteroatoms selected
 - O, S and N; R4 = aryl, heteroaryl, halo, etc.] which are mGluR5 modulators useful in the treatment or prevention of diseases and conditions in which mGluR5 is involved, including but not limited to psychiatric and mood disorders such as schizophrenia, anxiety, depression, bipolar disorders, and panic, as well as in the treatment of pain, Parkinson's disease, cognitive dysfunction, epilepsy, circadian rhythm and sleep disorders, such as shift-work induced sleep disorder and jet-lag, drug addiction, drug abuse, drug withdrawal, obesity and other diseases, were prepared Thus, amidation of pyridin-2-amine with 3-amino-5,6-diphenylpyrazine-2carboxylic acid afforded the amide II. The exemplified compds. I have mGluR5 inhibitory activity as shown by inhibition at 10 μM or less in the calcium flux assay or 100 μM or less or less in the PI assay. The invention is also directed to pharmaceutical compns. comprising compds. I. 863908-32-1P
- ΤТ RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES

(preparation of bipyridyl amides as modulators of metabotropic glutamate receptor-5)

863908-32-1 CAPLUS

CN Pyrazinecarboxamide, 3-amino-5,6-diphenyl-N-2-pyridinyl- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

from

RN

THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 6 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN 2005:493608 CAPLUS

ACCESSION NUMBER: DOCUMENT NUMBER: 143:43904

TITLE:

Preparation of pyrrolo[3,4-b]pyrazine-5,7(6H)-dione derivatives for treating obesity, psychiatric, and neurological disorders

INVENTOR(S): Cheng, Leifeng

PATENT ASSIGNEE(S): Astrazeneca AB, Swed.; Astrazeneca UK Limited SOURCE:

PCT Int. Appl., 26 pp.

LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

DOCUMENT TYPE:

GI

PATENT NO. KIND DATE APPLICATION NO. WO 2004-GB4934 WO 2005051953 A2 20050609 20041124 WO 2005051953 A3 20050728 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW RW: BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR, BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG AU 2004292493 A1 20050609 AU 2004-292493 20041124 CA 2004-2546318 CA 2546318 A1 20050609 20041124 EP 1701958 EP 2004-798641 A2 20060920 20041124 EP 1701958 В1 20070502 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, CY, TR, BG, CZ, EE, HU, PL, SK, HR, IS CN 1886405 20061227 CN 2004-80034802 А AT 2004-798641 AT 361301 Т 20070515 20041124 20070517 JP 2006-540602 JP 2007512298 Т 20041124 20071116 ES 2285544 Т3 ES 2004-4798641 20041124 IN 2006DN02621 20070824 IN 2006-DN2621 20060510 A US 2007099923 A1 20070503 US 2006-579830 20060517 HK 1096670 20071012 HK 2007-101236 20070201 A1 PRIORITY APPLN. INFO.: GB 2003-27331 A 20031125 WO 2004-GB4934 W 20041124 CASREACT 143:43904; MARPAT 143:43904 OTHER SOURCE(S):

OH

- AB The title compds. I [R1, R2 = Ph, thienyl, pyridyl, C1-C10-alkyl, C1-C10-alkyl, C2-C15-alkyl, C3-C15-cycloalkyl, R3 = C1-C15-alkyl, C3-C15-cycloalkyl, phenylC1-C4-alkyl, heteroaryl, heteroarylC1-C4-alkyl, R4(CH2)n, R4 = heterocycle, n = 0-4; X, Y = 0, S; Z = (0)n, n = 0, 1] were prepared and are designed to be used in the treatment of obesity, psychiatric disorders, neurol. disorders, immune, cardiovascular, reproductive, and endocrine disorders, septic shock, diseases related to respiratory and gastrointestinal systems, and extended abuse, addiction and/or relapse indications. As an example, 1,2-bis(4-chloropheyl) ethane-1,2-dione reacted with diaminomaleonitrile to give pyrazine-2,3-dicarbonitrile II which was treated with K0H/H2O2 in H2O, esterified, and hydrolyzed to give dicarboxylic acid III. III condensed with 4-FC6H4CRNL to give the mono-amide which cyclized to give the desired compound I (R1 = R2 = 4-C16H4, R3 = 4-FC6H4CR, X = Y = 0, Z = none).
- IT 811441-51-7P, 5,6-Bis(4-chlorophenyl)-3-[(piperidin-1ylamino)carbonyl]pyrazine-2-carboxylic acid 853578-19-5P 853578-23-1P
 - RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)
 - (preparation of pyrrolo[3,4-b]pyrazine-5,7(6H)-dione derivs. for treating obesity, psychiatric, neurol., immune, cardiovascular, reproductive, and endocrine disorders, septic shock, respiratory and gastrointestinal disorders)
- RN 811441-51-7 CAPLUS
- CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[(1piperidinylamino)carbonyl]- (9CI) (CA INDEX NAME)

- RN 853578-19-5 CAPLUS
- CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[[[(4-fluorophenyl)methyl]amino]carbonyl]- (9CI) (CA INDEX NAME)

RN 853578-23-1 CAPLUS

CN

Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[[(1,1-dimethylethyl)amino]carbonyl]- (9CI) (CA INDEX NAME)

L7 ANSWER 7 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:450934 CAPLUS

DOCUMENT NUMBER: 143:7731

TITLE: Preparation of pyrazine derivatives as adenosine receptor antagonists for treating neurological,

cardiovascular, and other diseases

INVENTOR(S): Yonishi, Satoshi; Aoki, Satoshi; Matsushima, Yuji; Akahane, Atsushi

PATENT ASSIGNEE(S): Fujisawa Pharmaceutical Co. Ltd., Japan

SOURCE: U.S. Pat. Appl. Publ., 37 pp.

CODEN: USXXCO
DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO. I	DATE
US 2005113387	A1	20050526	US 2004-972340 2	20041026
PRIORITY APPLN. INFO.:			EP 2003-905895 A 2	20031027
			EP 2004-902764 A 2	20040524
OTHER COURSE (C).	MADDAT	1/12.7721		

OTHER SOURCE(S): MARPAT 143:7731

AB Pyrazine derivative of formula I (with variables defined below) or salts thereof are claimed. The pyrazine compound I are adenosine antagonists and are useful for the prevention and/or treatment of depression, dementia (e.g. Alzheimer's disease, cerebrovascular dementia, dementia accompanying Parkinson's disease, etc.), Parkinson's disease, anxiety, pain,

cerebrovascular disease (e.g. stroke, etc.), heart failure and the like. A process for preparing the pyrazines and pharmaceutical compns. containing

them
are also claimed. For I, Rl is substituted pyridin-2-one or pyridine; R2
is H, OH, halogen, cyano, or optionally substituted lower alkyl, lower
alkenyl, lower alkynyl, lower alkoxy, aryloxy, arylthio, acyl, aryl,
heterocyclic group or amino; R3 and R4 are independently H, lower alkyl or
acyl; and R5 is optionally substituted lower alkyl, lower alkenyl, lower
alkynyl, cyano, aryl or heterocyclic group.

B 851087-21-3P, 3-Amino-6-(6-oxo-1,6-dihydro-3-pyridyl)-5-phenyl-2pyrazinecarboxamide 851087-22-4P, 3-Amino-6-(1-methyl-6-oxo-1,6dihydro-3-pyridyl)-5-phenyl-2-pyrazinecarboxamide 851087-39-3P,
3-Amino-5-(4-fiuorophenyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2pyrazinecarboxamide 851087-45-1P, 3-Amino-6-(1-isopropyl-6-oxo1,6-dihydro-3-pyridyl)-5-(4-methoxyphenyl)-2-pyrazinecarboxamide
851087-73-5P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)N-methoxy-N-methyl-3-phenyl-2-pyrazinecarboxamide 851087-95-1P,
3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2pyrazinecarboxamide
RL: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic
preparation); THU (Therapeutic use); BIOL (Biological study); PREP
(Preparation); RACT (Reactant or reagent); USES (Uses)

(drug candidate; preparation of pyrazine derivs. as adenosine receptor antagonists for treating neurol., cardiovascular, and other diseases) 851087-21-3 CAPLUS

CN Pyrazinecarboxamide, 3-amino-6-(1,6-dihydro-6-oxo-3-pyridinyl)-5-phenyl-(9CI) (CA INDEX NAME)

RN 851087-22-4 CAPLUS

CN Pyrazinecarboxamide, 3-amino-6-(1,6-dihydro-1-methyl-6-oxo-3-pyridinyl)-5phenyl- (9CI) (CA INDEX NAME)

$$\begin{array}{c} O \\ C-NH_2 \\ H_2N \\ N \\ N \\ N \\ \end{array}$$

RN 851087-39-3 CAPLUS

CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-5-(4-fluorophenyl)- (9CI) (CA INDEX NAME)

- RN 851087-45-1 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(4-methoxyphenyl)- (9CI) (CA INDEX NAME)

- RN 851087-73-5 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-N-methoxy-N-methyl-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-95-1 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-5-(2-pyridinyl)- (9CI) (CA INDEX NAME)

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851087-23-5P, 3-Amino-6-(1-ethyl-6-oxo-1,6-dihydro-3-pyridyl)-5-
phenyl-2-pyrazinecarboxamide 851087-24-6P, 3-Amino-6-(6-oxo-1-
propyl-1,6-dihydro-3-pyridyl)-5-phenyl-2-pyrazinecarboxamide
851087-25-7P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-
5-phenyl-2-pyrazinecarboxamide 851087-26-8P,
3-Amino-6-(6-isopropoxy-3-pyridyl)-5-phenyl-2-pyrazinecarboxamide
851087-36-0P, 3-Amino-6-(6-methoxy-3-pyridyl)-5-phenyl-2-
pyrazinecarboxamide 851087-37-1P, 3-Amino-5-(2-fluorophenyl)-6-
(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2-pyrazinecarboxamide
851087-38-2P, 3-Amino-5-(3-fluorophenyl)-6-(1-isopropyl-6-oxo-1,6-
dihydro-3-pyridyl)-2-pyrazinecarboxamide 851087-40-6P,
3-Amino-5-(2-chlorophenyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2-
pyrazinecarboxamide 851087-41-7P, 3-Amino-5-(3-chlorophenyl)-6-
(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2-pyrazinecarboxamide
851087-42-8P, 3-Amino-5-(4-chlorophenyl)-6-(1-isopropyl-6-oxo-1,6-
dihydro-3-pyridyl)-2-pyrazinecarboxamide 851087-43-9P,
3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-(2-methoxyphenyl)-2-
pyrazinecarboxamide 851087-44-0P, 3-Amino-6-(1-isopropyl-6-oxo-
1,6-dihydro-3-pyridyl)-5-(3-methoxyphenyl)-2-pyrazinecarboxamide
851087-46-2P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-
5-[2-(trifluoromethoxy)phenyl]-2-pyrazinecarboxamide 851087-47-3P
, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-[3-
(trifluoromethoxy)phenyl]-2-pyrazinecarboxamide 851087-48-4P.
3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-[4-
(trifluoromethoxy)phenyl]-2-pyrazinecarboxamide 851087-49-5P,
3-Amino-5-(3,4-difluorophenvl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-
2-pyrazinecarboxamide 851087-50-8P, 3-Amino-5-(3,5-
difluorophenyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2-
pvrazinecarboxamide 851087-51-9P, 3-Amino-5-(4-cyanophenyl)-6-(1-
isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2-pyrazinecarboxamide
851087-62-2P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-
N-methyl-5-phenyl-2-pyrazinecarboxamide 851087-63-3P,
3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-N,N-dimethyl-5-phenyl-
2-pyrazinecarboxamide 851087-65-5P, 3-Amino-6-(1-isopropyl-6-oxo-
1,6-dihydro-3-pyridyl)-5-phenyl-N-[(2-pyridyl)methyl]-2-
pyrazinecarboxamide 851087-66-6P, 3-Amino-N-(cyanomethyl)-6-(1-
isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-phenyl-2-pyrazinecarboxamide
851087-69-9P, 3-Amino-N-(2-hydroxyethyl)-6-(1-isopropyl-6-oxo-1,6-
dihydro-3-pyridyl)-5-phenyl-2-pyrazinecarboxamide 851087-70-2P.
3-Amino-N-cvclopropvl-6-(1-isopropvl-6-oxo-1,6-dihydro-3-pyridyl)-5-phenyl-
2-pyrazinecarboxamide 851087-77-9P, 3-Amino-N-[2-
(dimethylamino)ethyl]-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-phenyl-
2-pyrazinecarboxamide 851087-78-0P, 3-Amino-6-(1-isopropyl-6-oxo-
1,6-dihydro-3-pyridyl)-5-(2-methylphenyl)-2-pyrazinecarboxamide
851087-79-1P, 3-Amino-5-(2,3-difluorophenyl)-6-(1-isopropyl-6-oxo-
1,6-dihydro-3-pyridy1)-2-pyrazinecarboxamide 851087-80-4P,
3-Amino-5-(2,4-difluorophenyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-
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2-pyrazinecarboxamide 851087-81-5P, 3-Amino-5-(2,5-
difluorophenv1)-6-(1-isopropy1-6-oxo-1,6-dihydro-3-pyridy1)-2-
pyrazinecarboxamide 851087-82-6P, 3-Amino-5-(2-fury1)-6-(1-
isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2-pyrazinecarboxamide
851087-83-7P, 3-Amino-5-(3-furyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-
3-pyridy1)-2-pyrazinecarboxamide 851087-84-8P,
3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-(2-thienyl)-2-
pyrazinecarboxamide 851087-85-9P, 3-Amino-6-(1-isopropyl-6-oxo-
1,6-dihydro-3-pyridyl)-5-(3-thienyl)-2-pyrazinecarboxamide
851087-86-0P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-
5-(5-methyl-2-thienyl)-2-pyrazinecarboxamide 851087-87-1P.
3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-(1H-pyrazol-4-yl)-2-
pyrazinecarboxamide 851087-89-3P, 3-Amino-6-(1-isopropyl-6-oxo-
1,6-dihydro-3-pyridyl)-5-(3-pyridyl)-2-pyrazinecarboxamide
851087-90-6P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-
5-(4-pyridyl)-2-pyrazinecarboxamide 851087-91-7P,
3-Amino-5-(4-fluorophenyl)-6-(1-methyl-6-oxo-1,6-dihydro-3-pyridyl)-2-
pyrazinecarboxamide 851087-92-8P, 3-Amino-5-(2-fury1)-6-(1-
methyl-6-oxo-1,6-dihydro-3-pyridyl)-2-pyrazinecarboxamide
851087-93-9P, 3-Amino-6-(1-methyl-6-oxo-1,6-dihydro-3-pyridyl)-5-
(2-thienv1)-2-pyrazinecarboxamide 851088-51-2P,
3-Amino-5-(4-fluorophenvl)-6-(1-isopropvl-6-oxo-1,6-dihydro-3-pyridyl)-N-
methyl-2-pyrazinecarboxamide
RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU
(Therapeutic use); BIOL (Biological study); PREP (Preparation); USES
(Uses)
   (drug candidate; preparation of pyrazine derivs. as adenosine receptor
   antagonists for treating neurol., cardiovascular, and other diseases)
```

RN 851087-23-5 CAPLUS
CN Pyrazinecarboxamide, 3-amino-6-(1-ethyl-1,6-dihydro-6-oxo-3-pyridinyl)-5phenyl- (9C1) (CA INDEX NAME)

RN 851087-24-6 CAPLUS

CN Pyrazinecarboxamide, 3-amino-6-(1,6-dihydro-6-oxo-1-propyl-3-pyridinyl)-5phenyl- (9CI) (CA INDEX NAME)

- RN 851087-25-7 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-26-8 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[6-(1-methylethoxy)-3-pyridinyl]-5-phenyl-(9CI) (CA INDEX NAME)

- RN 851087-36-0 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-(6-methoxy-3-pyridinyl)-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-37-1 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(2-fluorophenyl)- (9CI) (CA INDEX NAME)

- RN 851087-38-2 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(3-fluorophenyl)- (9CI) (CA INDEX NAME)

- RN 851087-40-6 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(2-chlorophenyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

- RN 851087-41-7 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(3-chlorophenyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

- RN 851087-42-8 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(4-chlorophenyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

- RN 851087-43-9 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-5-(2-methoxyphenyl)- (9CI) (CA INDEX NAME)

- RN 851087-44-0 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(3-methoxyphenyl)- (9CI) (CA INDEX NAME)

- RN 851087-46-2 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-5-[2-(trifluoromethoxy)phenyl]- (9CI) (CA INDEX NAME)

- RN 851087-47-3 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-[3-(trifluoromethoxy)phenyl]- (9CI) (CA INDEX NAME)

- RN 851087-48-4 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-[4-(trifluoromethoxy)phenyl]- (9CI) (CA INDEX NAME)

RN 851087-49-5 CAPLUS

CN Pyrazinecarboxamide, 3-amino-5-(3,4-difluorophenyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

RN 851087-50-8 CAPLUS

CN Pyrazinecarboxamide, 3-amino-5-(3,5-difluorophenyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

RN 851087-51-9 CAPLUS

CN Pyrazinecarboxamide, 3-amino-5-(4-cyanopheny1)-6-[1,6-dihydro-1-(1methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

- RN 851087-62-2 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-N-methyl-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-63-3 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-N,N-dimethyl-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-65-5 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-phenyl-N-(2-pyridinylmethyl)- (9CI) (CA INDEX NAME)

- RN 851087-66-6 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-N-(cyanomethyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-69-9 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-N-(2-hydroxyethyl)-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-70-2 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-N-cyclopropyl-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-77-9 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-N-[2-(dimethylamino)ethyl]-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-78-0 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(2-methylphenyl)- (9CI) (CA INDEX NAME)

- RN 851087-79-1 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(2,3-difluorophenyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

- RN 851087-80-4 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(2,4-difluorophenyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

- RN 851087-81-5 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(2,5-difluorophenyl)-6-[1,6-dihydro-1-(1methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

- RN 851087-82-6 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(2-furanyl)- (9CI) (CA INDEX NAME)

- RN 851087-83-7 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-5-(3-furanyl)- (9CI) (CA INDEX NAME)

- RN 851087-84-8 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(2-thienyl)- (9CI) (CA INDEX NAME)

- RN 851087-85-9 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(3-thienyl)- (9CI) (CA INDEX NAME)

- RN 851087-86-0 CAPLUS CN
 - Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-5-(5-methyl-2-thienyl)- (9CI) (CA INDEX NAME)

- RN 851087-87-1 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-5-(1H-pyrazol-4-yl)- (9CI) (CA INDEX NAME)

- RN 851087-89-3 CAPLUS
- Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-CN pyridinyl]-5-(3-pyridinyl)- (9CI) (CA INDEX NAME)

- RN 851087-90-6 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-5-(4-pyridinyl)- (9CI) (CA INDEX NAME)

- RN 851087-91-7 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-(1,6-dihydro-1-methyl-6-oxo-3-pyridinyl)-5-(4-fluorophenyl)- (9CI) (CA INDEX NAME)

- RN 851087-92-8 CAPLUS
- CN Pyracinecarboxamide, 3-amino-6-(1,6-dihydro-1-methyl-6-oxo-3-pyridinyl)-5-(2-furanyl)- (9CI) (CA INDEX NAME)

- RN 851087-93-9 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-(1,6-dihydro-1-methyl-6-oxo-3-pyridinyl)-5-(2-thienyl) - (9CI) (CA INDEX NAME)

- RN 851088-51-2 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-5-(4-fluorophenyl)-N-methyl- (9CI) (CA INDEX NAME)

L7 ANSWER 8 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN 2005:395298 CAPLUS

ACCESSION NUMBER: DOCUMENT NUMBER: 142:447235

TITLE: Preparation of pyrazines as adenosine A1 and A2a receptor antagonists and their pharmaceutical compositions

INVENTOR(S): Yonishi, Satoshi; Aoki, Satoshi; Matsushima, Yuji;

Akahane, Atsushi Fujisawa Pharmaceutical Co., Ltd., Japan PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 152 pp.

CODEN: PIXXD2 Patent

DOCUMENT TYPE: LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT	NO.	KIN	KIND DATE			APPLICATION NO.						DATE						
WO 200	WO 2005040151					0506	WO 2004-JP16193						20041025					
W:	AE, A																	
	CN, C	CO, CF	, CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FΙ,	GB,	GD,			
	GE, (GH, GN	, HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	KZ,	LC,			
	LK, I	LR, LS	, LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	MZ,	NA,	NI,			
	NO, I	NZ, ON	, PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,			
	TJ,	IM, IN	, TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	ZW			
RW	: BW, 0	GH, GN	, KE,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,			
	AZ, I	BY, KG	, KZ,	MD,	RU,	TJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,			
	EE, I	ES, FI	, FR,	GB,	GR,	HU,	IE,	IT,	LU,	MC,	NL,	PL,	PT,	RO,	SE,			
	SI, S	SK, TF	, BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,			
	SN,	ID, TO																
AU 200	AU 2004283990					A1 20050506				AU 2004-283990								
									CA 2004-2543644									
EP 167	EP 1678160			A1 20060712			EP 2004-793294						20041025					
R:	AT, I	BE, CH	, DE,	DK,	ES,	FR,	GB,	GR,	IT,	LI,	LU,	NL,	SE,	MC,	PT,			
	IE, S	SI, FI	, RO,	CY,	TR,	BG,	CZ,	EE,	HU,	PL,	SK							
CN 187	CN 1871231					1129	CN 2004-80031570 BR 2004-15863 JP 2006-519017 MX 2006-PA4575						20041025					
BR 200	BR 2004015863					0109		BR 2	004-	1586	3		2	0041	025			
JP 200	JP 2007510620					0426		JP 2	006-	5190	17		2	0041	025			
MX 200	MX 2006PA04575					1120	MX 2006-PA4575						20060425					
NO 200	NO 2006002303					0719		NO 2	006-	2303			2	0060	522			
PRIORITY AP	PRIORITY APPLN. INFO.:							AU 2	003-	9058	95		A 2	0031	027			
								AU 2	004-	9027	64		A 2	0040	524			
								WO 2	004-	JP16	193	1	7 2	0041	025			
OTHER SOURC	E(S):		CAS	REAC	T 14	2:44	7235	; MA	RPAT	142	: 447	235						

$$R^{1}$$
 N
 R^{2}
 R^{3}
 R^{3}
 R^{4}
 R^{4}

II

- AB Title compound I (wherein Rl = N,3-disubstituted 2(1H)-pyridinonyl, 2-alkoxypyridinyl; R2 = H, OH, halo, CN, (un)substituted lower alk(en/yn)yl, alkoxy, aryloxy, arylthio, acyl, aryl, heterocyclyl or amino; R3, R4 = independently H, lower alkyl, acyl; and their salts) and their salts were prepared as adenosine receptor antagonists. For example, compound II was prepared by etherification of 5-(5-Amino-6-bromo-3-phenyl-pyrazinyl)-l-isopropyl-2(1H)-pyridinone (preparation given) with phenol. II showed binding to the human Al adenosine receptor with Ki = 1.57 nM and to the human A2a adenosine receptor with Ki = 0.32 nM. Thus, I are useful as Al receptor and A2a receptor dual antagonists and for the prevention and/or treatment of depression, dementia (e.g. Alzhelmer's disease, cerebrovascular dementia, dementia accompanying Parkinson's disease, etc.), Parkinson's disease, anxiety, pain, cerebrovascular disease (e.g. stroke, etc.), heart failure and the like (no data).
- Stroke, etc.), heart raillire and the like (No dazi) spridyl)-5-phenyl-2pyrazinecarboxamide 851087-39-3P, 3-Amino-5-(4-fluorophenyl)-6(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2-pyrazinecarboxamide
 851087-45-1P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)5-(4-methoxyphenyl)-2-pyrazinecarboxamide 851087-73-5P,
 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-N-methoxy-N-methyl-5phenyl-2-pyrazinecarboxamide 851087-95-1P, 3-Amino-6-(1isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-N-(2-pyridyl)-2-pyrazinecarboxamide
 Ri: PAC (Pharmacological activity); RCT (Reactant); SPN (Synthetic
 preparation); THU (Therapeutic use); BIOL (Biological study); PREP
 (Preparation); RACT (Reactant or reagent); USES (USES)
- (drug candidate; preparation of pyrazines as adenosine receptor antagonists)
- CN Pyrazinecarboxamide, 3-amino-6-(1,6-dihydro-6-oxo-3-pyridiny1)-5-phenyl-(9CI) (CA INDEX NAME)

- RN 851087-39-3 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(4-fluorophenyl)- (9CI) (CA INDEX NAME)

- RN 851087-45-1 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(4-methoxyphenyl)- (9CI) (CA INDEX NAME)

- RN 851087-73-5 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-N-methoxy-N-methyl-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-95-1 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-5-(2-pyridinyl)- (9CI) (CA INDEX NAME)

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851087-22-4P, 3-Amino-6-(1-methyl-6-oxo-1,6-dihydro-3-pyridyl)-5-
phenyl-2-pyrazinecarboxamide 851087-23-5P, 3-Amino-6-(1-ethyl-6-
oxo-1,6-dihydro-3-pyridyl)-5-phenyl-2-pyrazinecarboxamide
851087-24-6P, 3-Amino-6-(6-oxo-1-propyl-1,6-dihydro-3-pyridyl)-5-
phenyl-2-pyrazinecarboxamide 851087-25-7P, 3-Amino-6-(1-
isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-phenyl-2-pyrazinecarboxamide
851087-26-8P, 3-Amino-6-(6-isopropoxy-3-pyridyl)-5-phenyl-2-
pyrazinecarboxamide 851087-36-0P, 3-Amino-6-(6-methoxy-3-
pyridyl)-5-phenyl-2-pyrazinecarboxamide 851087-37-1P.
3-Amino-5-(2-fluorophenyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2-
pyrazinecarboxamide 851087-38-2P, 3-Amino-5-(3-fluorophenyl)-6-
(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2-pyrazinecarboxamide
851087-40-6P, 3-Amino-5-(2-chlorophenyl)-6-(1-isopropyl-6-oxo-1,6-
dihydro-3-pyridyl)-2-pyrazinecarboxamide 851087-41-7P,
3-Amino-5-(3-chlorophenyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2-
pyrazinecarboxamide 851087-42-8P, 3-Amino-5-(4-chlorophenyl)-6-
(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2-pyrazinecarboxamide
851087-43-9P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-
5-(2-methoxyphenyl)-2-pyrazinecarboxamide 851087-44-0P,
3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-(3-methoxyphenyl)-2-
pyrazinecarboxamide 851087-46-2P, 3-Amino-6-(1-isopropyl-6-oxo-
1,6-dihydro-3-pyridy1)-5-[2-(trifluoromethoxy)pheny1]-2-
pyrazinecarboxamide 851087-47-3P, 3-Amino-6-(1-isopropyl-6-oxo-
1,6-dihydro-3-pyridyl)-5-[3-(trifluoromethoxy)phenyl]-2-
pyrazinecarboxamide 851087-48-4P, 3-Amino-6-(1-isopropyl-6-oxo-
1,6-dihydro-3-pyridyl)-5-[4-(trifluoromethoxy)phenyl]-2-
pyrazinecarboxamide 851087-49-5P, 3-Amino-5-(3,4-difluorophenyl)-
6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2-pyrazinecarboxamide
851087-50-8P, 3-Amino-5-(3,5-difluorophenyl)-6-(1-isopropyl-6-oxo-
1,6-dihydro-3-pyridyl)-2-pyrazinecarboxamide 851087-51-9P,
3-Amino-5-(4-cvanophenyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2-
pyrazinecarboxamide 851087-62-2P, 3-Amino-6-(1-isopropyl-6-oxo-
1,6-dihydro-3-pyridyl)-N-methyl-5-phenyl-2-pyrazinecarboxamide
851087-63-3P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-
N, N-dimethyl-5-phenyl-2-pyrazinecarboxamide 851087-65-5P,
3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-phenyl-N-[(2-
pyridyl)methyl]-2-pyrazinecarboxamide 851087-66-6P,
3-Amino-N-(cvanomethyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-
phenyl-2-pyrazinecarboxamide 851087-69-9P, 3-Amino-N-(2-
hydroxyethyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-phenyl-2-
pyrazinecarboxamide 851087-70-2P, 3-Amino-N-cyclopropyl-6-(1-
isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-phenyl-2-pyrazinecarboxamide
851087-77-9P, 3-Amino-N-[2-(dimethylamino)ethyl]-6-(1-isopropyl-6-
oxo-1,6-dihydro-3-pyridy1)-5-pheny1-2-pyrazinecarboxamide
851087-78-0P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-
5-(2-methylphenyl)-2-pyrazinecarboxamide 851087-79-1P,
3-Amino-5-(2,3-difluorophenyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-
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2-pyrazinecarboxamide 851087-80-4P, 3-Amino-5-(2,4difluorophenv1)-6-(1-isopropy1-6-oxo-1,6-dihydro-3-pyridy1)-2pyrazinecarboxamide 851087-81-5P, 3-Amino-5-(2,5-difluorophenyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2-pyrazinecarboxamide 851087-82-6P, 3-Amino-5-(2-fury1)-6-(1-isopropy1-6-oxo-1,6-dihydro-3-pyridy1)-2-pyrazinecarboxamide 851087-83-7P, 3-Amino-5-(3-furyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-2pyrazinecarboxamide 851087-84-8P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-(2-thienyl)-2-pyrazinecarboxamide 851087-85-9P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-(3-thienv1)-2-pyrazinecarboxamide 851087-86-0P. 3-Amino-6-(1-isopropy1-6-oxo-1,6-dihydro-3-pyridy1)-5-(5-methy1-2-thieny1)-2-pyrazinecarboxamide 851087-87-1P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-(1H-pyrazol-4-yl)-2-pyrazinecarboxamide 851087-89-3P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-(3-pyridy1)-2-pyrazinecarboxamide 851087-90-6P, 3-Amino-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-5-(4-pyridyl)-2pyrazinecarboxamide 851087-91-7P, 3-Amino-5-(4-fluorophenyl)-6-(1-methyl-6-oxo-1,6-dihydro-3-pyridyl)-2-pyrazinecarboxamide 851087-92-8P, 3-Amino-5-(2-furyl)-6-(1-methyl-6-oxo-1,6-dihydro-3pvridv1)-2-pvrazinecarboxamide 851087-93-9P, 3-Amino-6-(1-methyl-6-oxo-1,6-dihydro-3-pyridyl)-5-(2-thienyl)-2pyrazinecarboxamide 851088-51-2P, 3-Amino-5-(4-fluorophenyl)-6-(1-isopropyl-6-oxo-1,6-dihydro-3-pyridyl)-N-methyl-2-pyrazinecarboxamide RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (drug candidate; preparation of pyrazines as adenosine receptor antagonists) 851087-22-4 CAPLUS

Pyrazinecarboxamide, 3-amino-6-(1,6-dihydro-1-methyl-6-oxo-3-pyridinyl)-5-

phenvl- (9CI) (CA INDEX NAME)

RN

CN

RN 851087-23-5 CAPLUS

CN Pyrazinecarboxamide, 3-amino-6-(1-ethyl-1,6-dihydro-6-oxo-3-pyridinyl)-5phenyl- (9CI) (CA INDEX NAME)

- RN 851087-24-6 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-(1,6-dihydro-6-oxo-1-propy1-3-pyridiny1)-5phenyl- (9CI) (CA INDEX NAME)

- RN 851087-25-7 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-26-8 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[6-(1-methylethoxy)-3-pyridinyl]-5-phenyl-(9CI) (CA INDEX NAME)

- RN 851087-36-0 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-(6-methoxy-3-pyridinyl)-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-37-1 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(2-fluorophenyl)- (9CI) (CA INDEX NAME)

- RN 851087-38-2 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(3-fluorophenyl)- (9CI) (CA INDEX NAME)

- RN 851087-40-6 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(2-chlorophenyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

- RN 851087-41-7 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(3-chloropheny1)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

- RN 851087-42-8 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(4-chlorophenyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

- RN 851087-43-9 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(2-methoxyphenyl)- (9CI) (CA INDEX NAME)

$$\begin{array}{c|c} & & & \\ &$$

- RN 851087-44-0 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(3-methoxyphenyl)- (9CI) (CA INDEX NAME)

- RN 851087-46-2 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-[2-(trifluoromethoxy)phenyl]- (9CI) (CA INDEX NAME)

- RN 851087-47-3 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-[3-(trifluoromethoxy)phenyl]- (9CI) (CA INDEX NAME)

- RN 851087-48-4 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-[4-(trifluoromethoxy)phenyl]- (9CI) (CA INDEX NAME)

- F3C-0
- RN 851087-49-5 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(3,4-difluorophenyl)-6-[1,6-dihydro-1-(1methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

- RN 851087-50-8 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(3,5-difluorophenyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

RN 851087-51-9 CAPLUS CN Pyrazinecarboxamide, 3-amino-5-(4-cyanophenyl)-6-[1,6-dihydro-1-(1methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

RN 851087-62-2 CAPLUS CN

Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-N-methyl-5-phenyl- (9CI) (CA INDEX NAME)

RN 851087-63-3 CAPLUS

Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-CN pyridinyl]-N, N-dimethyl-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-65-5 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-phenyl-N-(2-pyridinylmethyl)- (9CI) (CA INDEX NAME)

- RN 851087-66-6 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-N-(cyanomethyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-69-9 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-N-(2-hydroxyethyl)-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-70-2 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-N-cyclopropyl-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-77-9 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-N-[2-(dimethylamino)ethyl]-5-phenyl- (9CI) (CA INDEX NAME)

- RN 851087-78-0 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(2-methylphenyl)- (9CI) (CA INDEX NAME)

$$\begin{array}{c} \text{O} \\ \text{C} - \text{NH}_2 \\ \text{H}_2 \text{N} \\ \text{N} \\ \text{Me} \\ \end{array} \\ \begin{array}{c} \text{Pr-i} \\ \text{O} \\ \end{array}$$

- RN 851087-79-1 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(2,3-difluorophenyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

- RN 851087-80-4 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(2,4-difluorophenyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

- RN 851087-81-5 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5-(2,5-difluorophenyl)-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]- (9CI) (CA INDEX NAME)

- RN 851087-82-6 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-5-(2-furanyl)- (9CI) (CA INDEX NAME)

- RN 851087-83-7 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(3-furanyl)- (9CI) (CA INDEX NAME)

- RN 851087-84-8 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(2-thienyl)- (9CI) (CA INDEX NAME)

- RN 851087-85-9 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(3-thienyl)- (9CI) (CA INDEX NAME)

- RN 851087-86-0 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl]-5-(5-methyl-2-thienyl)- (9CI) (CA INDEX NAME)

- RN 851087-87-1 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(1H-pyrazol-4-yl)- (9CI) (CA INDEX NAME)

RN 851087-89-3 CAPLUS
CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3pyridinyl)-5-(3-pyridinyl)- (9CI) (CA INDEX NAME)

- RN 851087-90-6 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(4-pyridinyl)- (9CI) (CA INDEX NAME)

- RN 851087-91-7 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-(1,6-dihydro-1-methyl-6-oxo-3-pyridinyl)-5-(4-fluorophenyl)- (9CI) (CA INDEX NAME)

- RN 851087-92-8 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-(1,6-dihydro-1-methyl-6-oxo-3-pyridinyl)-5-(2-furanyl)- (9CI) (CA INDEX NAME)

- RN 851087-93-9 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-(1,6-dihydro-1-methyl-6-oxo-3-pyridinyl)-5-(2-thienyl)-(9CI) (CA INDEX NAME)

- RN 851088-51-2 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-6-[1,6-dihydro-1-(1-methylethyl)-6-oxo-3-pyridinyl]-5-(4-fluorophenyl)-N-methyl- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

THERE ARE 9 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

9 L7 ANSWER 9 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

2004:1127371 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 142:56364

TITLE: Preparation of 2,3-substituted 5,6-diaryl-pyrazine

derivatives as CB1 modulators INVENTOR(S): Cheng, Leifeng; Wilstermann, Michael

PATENT ASSIGNEE(S): Astrazeneca Ab, Swed. SOURCE: PCT Int. Appl., 54 pp.

CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

	PATENT NO.						KIND DATE					ICAT							
							A1 20041223			WO 2004-SE968									
		W:															CA, GB,		
																	KZ,		
																	NA,		
																	SL,		
		RW:															ZM,		
																	DE,		
																	RO,		
						BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	
	SN, TD, TG							2004	1223	AU 2004-247614						20040616			
										CA 2004-2527037									
	EP 1638956					A1 20060329			EP 2004-749010						20040616				
		R:													NL,	SE,	MC,	PT,	
												HU,							
	JP 2006527769							2006	1207	JP 2006-517042						20040616			
US 2007093505						A1 20070426				US 2005-561033									
PRIORITY APPLN. INFO.:										GB 2	003-	1426	1		A 2	20030	619		
							WO 2	004-	SE96	8		W 2	20040	616					
OTHER	OTHER COHECE/C).						DAT	1/2.	5626	4									

OTHER SOURCE(S): MARPAT 142:56364

GI

Title compds. I [wherein R1, R2 = independently (un) substituted Ph, AB thienvl, pyridinyl; R3, R4 = (CH2)nCO2R7, CH2OCH2R8, (CH2)gR9 with proviso, (un)substituted alkyl, etc.; R7 = (un)substituted cycloalkyl/cyclo/alkyl, (CH2)aphenyl, (un)saturated heterocyclyl; a = 0-4; R8 = (un)substituted alkyl, Ph, (un)saturated aromatic heterocyclyl; n = 0-4; q = 0-4; R9 = (un)substituted cycloalkyl, ph, aromatic heterocyclyl, saturated or partially unsatd. 5-12-membered heterocyclyl; and pharmaceutically acceptable salts thereof] were prepared as cannabinoid 1 (CB1) receptor modulators. Thus, reacting (DL)-alaninol with 5,6-Bis(4-chlorophenyl)-3-(tert-butoxycarbonyl)pyrazine-2-carboxylic acid (preparation given), followed by cyclization gave pyrazine II. I are active at the CB1 receptor (IC50 < 1 µM), most preferred compds. have IC50 < 200 nM. For instance, II exhibited an IC50 (hCB1) = 1.8 nM. Thus, I and their pharmaceutical compns. are useful for the treatment of obesity, psychiatric and neurol. disorders (no data).

II 811436-87-0P, 5,6-Bis(4-chlorophenyl)-3-[N-(2-hydroxy-1,1-dimethylethyl)carbamoyl]pyrazine-2-carboxylic acid tert-butyl ester 811436-90-5P, 5,6-Bis(4-chlorophenyl)-3-[N-(1-(hydroxymethyl)cyclophentyl)carbamoyl]pyrazine-2-carboxylic acid tert-butyl ester 811436-92-7P, 5,6-Bis(4-chlorophenyl)-3-[N-(2-hydroxy-1-methylethyl)carbamoyl]pyrazine-2-carboxylic acid tert-butyl ester 811436-95-0P, 5,6-Bis(4-chlorophenyl)-3-[N-(2-hydroxy-1-phenylethyl)carbamoyl]pyrazine-2-carboxylic acid tert-butyl ester 811436-99-3P, 5,6-Bis(4-chlorophenyl)-3-[N-(2-hydroxy-2-phenylethyl)carbamoyl]pyrazine-2-carboxylic acid tert-butyl ester RL: RCT (Reactant) spr (Synthetic preparation); PREP (Preparation); RACT (Reactant) or reagent)

(intermediate; preparation of 2,3-substituted 5,6-diaryl-pyrazines as CB1 modulators)

RN 811436-87-0 CAPLUS

CN Pyrazinecarboxylic acid, 5,6-bis(4-chloropheny1)-3-[[(2-hydroxy-1,1-dimethylethy1)amino]carbony1]-, 1,1-dimethylethy1 ester (9CI) (CA INDEX NAME)

- RN 811436-90-5 CAPLUS

- RN 811436-92-7 CAPLUS
- CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[[(2-hydroxy-1-methylethyl)amino]carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

- RN 811436-95-0 CAPLUS
- CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[[(2-hydroxy-1-

phenylethyl)amino]carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

RN 811436-98-3 CAPLUS

CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[[(2-hydroxy-2-phenylethyl)amino]carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 10 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2004:1127370 CAPLUS

DOCUMENT NUMBER: 142:56363

TITLE: Preparation of 5,6-bis(4-chlorophenyl)-N-piperidin-1yl-3-(piperidin-1-ylcarbonyl)pyrazine-2-carboxamide

for treatment of obesity

INVENTOR(S): Cheng, Leifeng

PATENT ASSIGNEE(S): Astrazeneca Ab, Swed. SOURCE: PCT Int. Appl., 24 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent

LANGUAGE: English
FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

PRIORITY APPLN. INFO.: GB 2003-14049 A 20030618 GI

AB 5,6-Bis(4-chlorophenyl)-N-piperidin-1-yl-3-(piperidin-1-yl-acarbonyl)pyrazine-2-carboxamide (I) was prepared by reacting 4-ClC6H4CH0 with NaCN/EtOH which gave 1,2-bis(4-chlorophenyl)-2-hydroxyethanone (II). II was oxidized to the ethane-1,2-dione which was condensed with diaminomaleonitrile to give pyrazine III. III was converted to the corresponding 2,3-dicarboxylic acid which was treated with AcCl to give furo[3,4-b]pyrazine-5,7-dione IV. IV was then subsequently reacted with piperidine/MecN and oxalyl chloride/1-piperidinamine/CH2Cl2 to give the title compound that is intended to be used to treat obesity, psychiatric and neurol disorders.

IT 810685-52-0P

for

RN

CN

RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(preparation of bis(chlorophenyl)piperidinylpyrazinecarboxamide derivative

treating obesity, psychiatric disorders, and neurol. disorders) 810685-52-0 CAPLUS

Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-1-piperidinyl-3-(1-piperidinylcarbonyl)- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS
RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 11 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2004:1127366 CAPLUS

DOCUMENT NUMBER: 142:56362

TITLE: Preparation of 3-substituted 5,6-diaryl-pyrazine-2-

carboxamide and 2-sulfonamide derivatives as

cannabinoid receptor 1 (CB1) modulators INVENTOR(S): Cheng, Leifeng

INVENTOR(S): Cheng, Leifeng
PATENT ASSIGNEE(S): Astrazeneca AB, Swed.

SOURCE: PCT Int. Appl., 120 pp.

CODEN: PIXXD2

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT: 1 PATENT INFORMATION:

				KIND DATE			APPLICATION NO.											
							WO 2004-SE970											
	W:	ΑE,	AG,	AL,	AM,	AT,	AU,	AZ,	BA,	BB,	BG,	BR,	BW,	BY,	BZ,	CA,	CH,	
		CN,	CO,	CR,	CU,	CZ,	DE,	DK,	DM,	DZ,	EC,	EE,	EG,	ES,	FI,	GB,	GD,	
		GE,	GH,	GM,	HR,	HU,	ID,	IL,	IN,	IS,	JP,	KE,	KG,	KP,	KR,	ΚZ,	LC,	
		LK,	LR,	LS,	LT,	LU,	LV,	MA,	MD,	MG,	MK,	MN,	MW,	MX,	ΜZ,	NA,	NI,	
		NO,	NZ,	OM,	PG,	PH,	PL,	PT,	RO,	RU,	SC,	SD,	SE,	SG,	SK,	SL,	SY,	
		ТJ,	TM,	TN,	TR,	TT,	TZ,	UA,	UG,	US,	UZ,	VC,	VN,	YU,	ZA,	ZM,	zw	
	RW:	BW,	GH,	GM,	KΕ,	LS,	MW,	MZ,	NA,	SD,	SL,	SZ,	TZ,	UG,	ZM,	ZW,	AM,	
		AZ,	BY,	KG,	KZ,	MD,	RU,	ТJ,	TM,	AT,	BE,	BG,	CH,	CY,	CZ,	DE,	DK,	
		EE,	ES,	FΙ,	FR,	GB,	GR,	HU,	ΙE,	IT,	LU,	MC,	NL,	PL,	PT,	RO,	SE,	
		SI,	SK,	TR,	BF,	ВJ,	CF,	CG,	CI,	CM,	GA,	GN,	GQ,	GW,	ML,	MR,	NE,	
		SN,	TD,	TG														
ΑU	2004247616			A1 20041223			AU 2004-247616											
										CA 2004-2527035								
EΡ	1638	953			A1		20060329 EP 2004-749012					20040616						
	R:	ΑT,																
								MK,										
BR	2004011508				A		20060725			BR 2004-11508					20040616			
										CN 2004-80017200								
JP	P 2006527771				T		20061207			JP 2006-517044				20040616				

NO 2005005919	A	20060216	NO	2005-5919		20051213
MX 2005PA13711	A	20060308	MX	2005-PA13711		20051215
US 2007093484	A1	20070426	US	2005-560862		20051215
PRIORITY APPLN. INFO.:			GB	2003-14057	A	20030618
			WO	2004-SE970	W	20040616

OTHER SOURCE(S): MARPAT 142:56362

AR Title compds. I [wherein R1, R2 = independently (un)substituted Ph, thienyl, pyridinyl; R3 = X-Y-NR5R6; X = absent, CO, or SO2; Y = absent, NH optionally substituted by an alkyl group; R5, R6 = independently (un) substituted amino/alkyl, (CH2)r(phenyl)s, (un) saturated 5-8-membered heterocyclyl; R5 = H and R6 = defined above; or R5NR6 = (un)substituted (un) saturated 5-8-membered heterocyclyl; r = 0-4; s = 1 when r = 0, otherwise s = 1 or 2; R5NR6 = (un)substituted (un)saturated 5-8-membered heterocyclyl; R4 = (CH2) nCO2R7; n = 0-4; R7 = (un) substituted cycloalkyl/cyclo/alkyl, (CH2) nphenyl, saturated or partially unsatd. 5-8-membered heterocyclyl, CONH2 and derivs.; n = defined as above; and pharmaceutically acceptable salts thereof] were prepared as cannabinoid 1 (CB1) receptor modulators. For example, reacting 3-(tert-butoxycarbonyl)-5,6-bis(4-chlorophenyl)pyrazine-2-carboxylic acid (preparation given) with tert-butylhydrazine hydrochloride gave pyrazine II. I are active at the CB1 receptor (IC50 < 1 µM), most preferred compds. have IC50 < 200 nM. For instance, II exhibited an IC50 (hCB1) = 1.8 nM. Thus, I and their pharmaceutical compns. are useful for the treatment of obesity, psychiatric and neurol. disorders (no data).

ΙI

IT 811441-12-OP, 5,6-Bis(4-chlorophenyl)-3-(cyanomethyl)-N-(piperidin1-yl)pyrazine-2-carboxamide 811441-34-6P, tert-Butyl
[[1-[15,6-bis(4-chlorophenyl)-3-[[(piperidin-1-yl)amino]carbonyl]pyrazin-2yl]methyl]-1H-1,2,3-triazol-4-yl]methyl]-carbomate 811441-35-7P
RL: PRC (Pharmacological activity); RCT (Reactant); SPN (Synthetic
preparation); THU (Therapeutic use); BIOL (Biological study); PREP
(Preparation); RACT (Reactant or reagent); USES (Uses)

(drug candidate; preparation of 3-substituted 5,6-diarylpyrazine-2-carboxamide and 2-sulfonamide derivs. as CB1 modulators)

RN 811441-12-0 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chloropheny1)-3-(cyanomethy1)-N-1piperidiny1- (9CI) (CA INDEX NAME)

- RN 811441-34-6 CAPLUS
- CN Carbamic acid, [[1-[6,6-bis(4-chlorophenyl)-3-[(1-piperidinylamino)carbonyl]pyrazinyl]methyl]-lH-1,2,3-triazol-4-yl]methyl]-,1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

- RN 811441-35-7 CAPLUS
- CN Carbamic acid, [[1-[5,6-bis(4-chlorophenyl)-3-[(1-piperidinylamino)carbonyl)pyrazinyl]methyl]-1H-1,2,3-triazol-5-yl]methyl]-1,1-dimethylethyl ester (901) (CA INDEX NAME)

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811436-92-7P, tert-Butvl 5,6-bis(4-chlorophenvl)-3-[[(2-hvdroxv-1-
methylethyl)amino]carbonyl]pyrazine-2-carboxylate 811440-95-6P,
tert-Butyl 5,6-bis(4-chlorophenyl)-3-[[(piperidin-1-
vl)amino]carbonyl]pyrazine-2-carboxylate 811440-96-7P, Butyl
5,6-bis(4-chlorophenyl)-3-[[(piperidin-1-yl)amino]carbonyl]pyrazine-2-
carboxylate 811440-97-8P, Cyclohexyl 5,6-bis(4-chlorophenyl)-3-
[[(piperidin-1-v1)amino]carbonv1]pvrazine-2-carboxvlate
811440-98-9P, Benzyl 5,6-bis(4-chlorophenyl)-3-[[(piperidin-1-
yl)amino]carbonyl]pyrazine-2-carboxylate 811440-99-0P,
tert-Butyl 5,6-bis(4-chlorophenyl)-3-[[(cis-2-
hydroxycvclohexyl)amino]carbonyl]pyrazine-2-carboxylate
811441-00-6P, tert-Butyl 5,6-bis(4-chlorophenyl)-3-[[(trans-2-
hydroxycyclohexyl)amino]carbonyl]pyrazine-2-carboxylate
811441-01-7P, tert-Butyl 5,6-bis(4-chlorophenyl)-3-[[2-[4-
(trifluoromethyl)phenyl]hydrazino]carbonyl]pyrazine-2-carboxvlate
811441-02-8P, tert-Butyl 5,6-bis(4-chlorophenyl)-3-[[(morpholin-4-
yl)amino]carbonyl]pyrazine-2-carboxylate 811441-03-9P,
tert-Butvl 5,6-bis(4-chlorophenvl)-3-[[2-(tert-
butv1)hvdrazino|carbonv1|pvrazine-2-carboxvlate 811441-04-0P,
3-(tert-Butoxymethyl)-5,6-bis(4-chlorophenyl)-N-(piperidin-1-yl)pyrazine-2-
carboxamide 811441-08-4P, 5,6-Bis(4-chlorophenyl)-3-
[(cyclohexylidene)methyl]-N-(piperidin-1-yl)pyrazine-2-carboxamide
811441-17-5P, 5,6-Bis(4-chlorophenyl)-3-(1-methoxyethyl)-N-
(piperidin-1-vl)pyrazine-2-carboxamide 811441-22-2P, tert-Butyl
5,6-bis(4-chlorophenyl)-3-[[(4,4-difluorocyclohexyl)amino]carbonyl]pyrazin
e-2-carboxylate 811441-23-3P, tert-Butyl 5,6-bis(4-chlorophenyl)-
3-[(pentylamino)carbonyl]pyrazine-2-carboxylate 811441-24-4P,
tert-Butyl 5,6-bis(4-chlorophenyl)-3-[[(1-ethylpropyl)amino]carbonyl]pyraz
ine-2-carboxylate 811441-25-5P, tert-Butyl 5,6-bis(4-
chlorophenyl)-3-[[(4,4-difluoropiperidin-1-yl)amino]carbonyl]pyrazine-2-
carboxylate 811441-27-7P, 5,6-Bis(4-chlorophenyl)-N-(piperidin-1-
y1)-3-[(4-propy1-1H-1,2,3-triazol-1-y1)methy1]pyrazine-2-carboxamide
811441-32-4P, 5,6-Bis(4-chlorophenyl)-3-[[5-(1-hydroxyethyl)-1H-
1,2,3-triazol-1-y1]methyl]-N-(piperidin-1-y1)pyrazine-2-carboxamide
811441-36-8P, 3-[[4-(Aminomethyl)-1H-1,2,3-triazol-1-yl]methyl]-
5,6-bis(4-chlorophenyl)-N-(piperidin-1-yl)pyrazine-2-carboxamide
hydrochloride 811441-37-9P, 3-[[5-(Aminomethy1)-1H-1,2,3-triazol-
1-y1]methy1]-5,6-bis(4-chloropheny1)-N-(piperidin-1-y1)pyrazine-2-
carboxamide hydrochloride 811441-38-0P, 5,6-Bis(4-chloropheny1)-
3-(phenoxymethyl)-N-(piperidin-1-yl)pyrazine-2-carboxamide
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811441-40-4P, 5,6-Bis(4-chlorophenyl)-3-[(morpholin-4-yl)methyl]-N-
(piperidin-1-v1)pyrazine-2-carboxamide 811441-42-6P,
5,6-Bis(4-chlorophenv1)-3-[(piperidin-1-v1)methv1]-N-(piperidin-1-
yl)pyrazine-2-carboxamide 811441-44-8P, 5,6-Bis(4-chlorophenyl)-
3-[(cyclohex-2-en-1-y1)oxy]methy1]-N-(piperidin-1-y1)pyrazine-2-
carboxamide 811441-47-1P, 5,6-Bis(4-chlorophenyl)-3-
[(cyclohexyloxy)methyl]-N-(piperidin-1-yl)pyrazine-2-carboxamide
811441-50-6P, 5,6-Bis(4-chlorophenyl)-N-(2-hydroxyethyl)-N'-
(piperidin-1-v1)pvrazine-2,3-dicarboxamide 811441-52-8P,
5,6-Bis(4-chlorophenvl)-N-(3-hvdroxybutyl)-N'-(piperidin-1-vl)pyrazine-2,3-
dicarboxamide 811441-53-9P, 5,6-Bis(4-chlorophenyl)-N-(3-
hydroxypropyl)-N'-(piperidin-1-yl)pyrazine-2,3-dicarboxamide
811441-54-0P, tert-Butyl 5,6-bis(4-methylphenyl)-3-[[(piperidin-1-
yl)amino]carbonyl]pyrazine-2-carboxvlate 811441-58-4P.
5,6-Bis(4-methylphenyl)-N-(piperidin-1-yl)-3-[(1H-tetrazol-1-
yl)methyl]pyrazine-2-carboxamide 811441-62-0P,
5,6-Bis(4-methylphenyl)-N-(piperidin-1-yl)-3-[(2H-tetrazol-2-
yl)methyl]pyrazine-2-carboxamide 811441-64-2P,
5,6-Bis(4-chlorophenyl)-N-(piperidin-1-yl)-3-[(2H-tetrazol-2-
vl)methyl]pyrazine-2-carboxamide 811441-65-3P,
5,6-Bis(4-chlorophenyl)-N-(piperidin-1-yl)-3-[(1H-tetrazol-1-
v1)methvllpvrazine-2-carboxamide 811441-66-4P,
5,6-Bis(4-chlorophenyl)-N-(4,4-difluorocyclohexyl)-3-[(2H-tetrazol-2-
v1)methv11pvrazine-2-carboxamide 811441-67-5P,
5,6-Bis(4-chlorophenyl)-N-(4,4-difluoropiperidin-1-yl)-3-[(2H-tetrazol-2-
vl)methyl]pyrazine-2-carboxamide 811441-68-6P,
5,6-Bis(4-chlorophenyl)-3-[(2-methoxyethoxy)methyl]-N-(piperidin-1-
yl)pyrazine-2-carboxamide 811441-71-1P, 5,6-Bis(4-chlorophenyl)-
3-((5-cvclopropvl-2H-tetrazol-2-vl)methvl]-N-(piperidin-1-vl)pvrazine-2-
carboxamide 811441-74-4P, 5,6-Bis(4-chlorophenyl)-3-[(5-
cyclopropyl-1H-tetrazol-1-yl)methyl]-N-(piperidin-1-yl)pyrazine-2-
carboxamide 811441-75-5P, 5,6-Bis(4-chlorophenyl)-3-[(5-methyl-
2H-tetrazo1-2-y1)methy1]-N-(piperidin-1-y1)pyrazine-2-carboxamide
811441-78-8P, 5,6-Bis(4-chlorophenyl)-3-[(5-methyl-1H-tetrazol-1-
yl)methyl]-N-(piperidin-1-yl)pyrazine-2-carboxamide 811441-79-9P
, tert-Butyl 6-(4-chlorophenyl)-5-(4-methylphenyl)-3-[[(piperidin-1-
yl)amino]carbonyl]pyrazine-2-carboxylate 811441-86-8P,
tert-Butyl 5-(4-chlorophenyl)-6-(4-methylphenyl)-3-[[(piperidin-1-
yl)amino]carbonyl]pyrazine-2-carboxylate 811441-87-9P,
6-(4-Chlorophenyl)-5-(4-methylphenyl)-N-(piperidin-1-yl)-3-[(2H-tetrazol-2-
v1)methv11pvrazine-2-carboxamide 811441-94-8P,
5-(4-Chlorophenyl)-6-(4-methylphenyl)-N-(piperidin-1-yl)-3-[(2H-tetrazol-2-
v1)methv1]pvrazine-2-carboxamide 811441-97-1P, tert-Butv1
5,6-bis(4-chlorophenyl)-3-[[(2-hydroxyethyl)(methyl)amino]carbonyl]pyrazin
e-2-carboxylate 811441-98-2P, 5,6-Bis(4-chlorophenyl)-3-
propoxypyrazine-2-carboxylic acid N-(piperidin-1-v1)amide
811442-03-2P, 5,6-Bis(4-chlorophenyl)-N-(piperidin-1-yl)-3-[(2H-
tetrazol-5-vl)methvl]pvrazine-2-carboxamide 811442-07-6P,
5,6-Bis(4-chlorophenyl)-3-[[5-(morpholin-4-yl)-2H-tetrazol-2-yl]methyl]-N-
(piperidin-1-yl)pyrazine-2-carboxamide 811442-08-7P,
5,6-Bis(4-chlorophenyl)-3-[[5-(morpholin-4-yl)-1H-tetrazol-1-yl]methyl]-N-
(piperidin-1-vl)pyrazine-2-carboxamide 811442-10-1P,
5,6-Bis(4-chlorophenyl)-N-(piperidin-1-v1)-3-[[5-(pyrrolidin-1-v1)-2H-
tetrazol-2-vl|methvl|pvrazine-2-carboxamide 811442-11-2P.
5,6-Bis(4-chlorophenyl)-N-(piperidin-1-yl)-3-[[5-(pyrrolidin-1-yl)-1H-
tetrazol-1-y1]methyl]pyrazine-2-carboxamide 811442-12-3P,
5,6-Bis(4-chlorophenyl)-3-[[5-(methylthio)-2H-tetrazol-2-yl]methyl]-N-
(piperidin-1-v1)pvrazine-2-carboxamide 811442-13-4P,
5,6-Bis(4-chloropheny1)-3-[[5-(methylthio)-1H-tetrazol-1-y1]methyl]-N-
(piperidin-1-yl)pyrazine-2-carboxamide 811442-14-5P,
5,6-Bis(4-chlorophenyl)-N-(4,4-difluorocyclohexyl)-3-
(methoxymethyl)pyrazine-2-carboxamide 811442-16-7P,
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5,6-Bis(4-chlorophenyl)-N-(4,4-difluorocyclohexyl)-3-[[(4fluorobenzyl)oxy|methyl|pyrazine-2-carboxamide 811442-19-0P, 5,6-Bis(4-chlorophenyl)-3-[(4,4-difluoropiperidin-1-yl)methyl]-N-(piperidin-1-yl)pyrazine-2-carboxamide 811442-21-4P, 5,6-Bis(4-chlorophenyl)-N-(4,4-difluorocyclohexyl)-3-[(4,4difluoropiperidin-1-yl)methyl]pyrazine-2-carboxamide 811442-22-5P , 5,6-Bis(4-chlorophenyl)-N-(4,4-difluoropiperidin-1-yl)-3-(methoxymethyl)pyrazine-2-carboxamide 811442-24-7P, 5,6-Bis(4-chlorophenv1)-3-[[4-(1-hvdroxyethv1)-1H-1,2,3-triazol-1vllmethvll-N-(piperidin-1-vl)pvrazine-2-carboxamide 811442-25-8P , 3-[[4-(Aminomethyl)-1H-1,2,3-triazol-1-yl]methyl]-5,6-bis(4chlorophenyl)-N-(piperidin-1-yl)pyrazine-2-carboxamide 811442-26-9P, 3-[[5-(Aminomethyl)-1H-1,2,3-triazol-1-yl]methyl]-5,6-bis(4-chlorophenyl)-N-(piperidin-1-yl)pyrazine-2-carboxamide RL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

(drug candidate; preparation of 3-substituted 5,6-diarylpyrazine-2-carboxamide and 2-sulfonamide derivs. as CB1 modulators)

RN 811436-92-7 CAPLUS

CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[[(2-hydroxy-1-methylethyl)amino]carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

RN 811440-95-6 CAPLUS

CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[(1piperidinylamino)carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

- RN 811440-96-7 CAPLUS
- CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[(1-piperidinylamino)carbonyl]-, butyl ester (9CI) (CA INDEX NAME)

- RN 811440-97-8 CAPLUS
- CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[(1piperidinylamino)carbonyl]-, cyclohexyl ester (9CI) (CA INDEX NAME)

- RN 811440-98-9 CAPLUS
- CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[(1-piperidinylamino)carbonyl]-, phenylmethyl ester (9CI) (CA INDEX NAME)

RN 811440-99-0 CAPLUS

CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[[[(1R,2S)-2-hydroxycyclohexyl]amino]carbonyl]-, 1,1-dimethylethyl ester, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

RN 811441-00-6 CAPLUS

CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[[[(1R,2R)-2-hydroxycyclohexyl]amino]carbonyl]-, 1,1-dimethylethyl ester, rel- (9CI) (CA INDEX NAME)

Relative stereochemistry.

- RN 811441-01-7 CAPLUS
- CN 2,3-Pyrazinedicarboxylic acid, 5,6-bis(4-chlorophenyl)-, mono(1,1-dimethylethyl) ester, 2-[4-(trifluoromethyl)phenyl]hydrazide (9C1) (CA INDEX NAME)

- RN 811441-02-8 CAPLUS
- CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[(4-morphollinylamino)carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

- RN 811441-03-9 CAPLUS
- 23.3-Pyrazinedicarboxylic acid, 5,6-bis(4-chlorophenyl)-, mono(1,1-dimethylethyl) ester, 2-(1,1-dimethylethyl)hydrazide (9CI) (CA INDEX NAME)

- RN 811441-04-0 CAPLUS
- CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-[(1,1-dimethylethoxy)methyl]-N-1-piperidinyl- (9CI) (CA INDEX NAME)

- RN 811441-08-4 CAPLUS
- CN Pyrazinecarboxamide, 5,6-bis(4-chloropheny1)-3-(cyclohexylidenemethyl)-N-1piperidinyl- (9CI) (CA INDEX NAME)

- RN 811441-17-5 CAPLUS
- CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-(1-methoxyethyl)-N-1-piperidinyl- (9CI) (CA INDEX NAME)

- RN 811441-22-2 CAPLUS
- CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[[(4,4-difluorocyclohexyl)amino]carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

- RN 811441-23-3 CAPLUS
- CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[(pentylamino)carbonyl], 1,1-dimethylethyl ester (9Cl) (CA INDEX NAME)

- RN 811441-24-4 CAPLUS
- CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[[(1ethylpropyl)amino]carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

- RN 811441-25-5 CAPLUS
- CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[[(4,4-difluoro-1-piperidinyl)amino]carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

- RN 811441-27-7 CAPLUS
- CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-1-piperidinyl-3-[(4-propyl-1H-1,2,3-triazol-1-yl)methyl]- (9CI) (CA INDEX NAME)

- RN 811441-32-4 CAPLUS
- CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-[[5-(1-hydroxyethyl)-1H-1,2,3-triazol-1-yl]methyl]-N-1-piperidinyl- (9CI) (CA INDEX NAME)

RN 811441-36-8 CAPLUS
CN Pyrazinecarboxamide, 3-[[4-(aminomethyl)-1H-1,2,3-triazol-1-yl]methyl]-5,6bis(4-chlorophenyl)-N-1-piperidinyl-, hydrochloride (9CI) (CA INDEX NAME)

RN 811441-37-9 CAPLUS
CN Pyrazinecarboxamide, 3-[[5-(aminomethyl)-1H-1,2,3-triazol-1-yl]methyl]-5,6bis(4-chlorophenyl)-N-1-piperidinyl-, hydrochloride (9C1) (CA INDEX NAME)

811441-38-0 CAPLUS RN CN

Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-(phenoxymethyl)-N-1-piperidinyl- (9CI) (CA INDEX NAME)

811441-40-4 CAPLUS RN

CN

RN 811441-42-6 CAPLUS
CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-1-piperidinyl-3-(1-piperidinylmethyl)- (9CI) (CA INDEX NAME)

- RN 811441-44-8 CAPLUS
- CN Pyrazinecarboxamide, 5,6-bis(4-chloropheny1)-3-[(2-cyclohexen-1-yloxy)methy1]-N-1-piperidiny1- (9CI) (CA INDEX NAME)

- RN 811441-47-1 CAPLUS
- CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-[(cyclohexyloxy)methyl]-N-1-piperidinyl- (9CI) (CA INDEX NAME)

- RN 811441-50-6 CAPLUS
- CN 2,3-Pyrazinedicarboxamide, 5,6-bis(4-chlorophenyl)-N-(2-hydroxyethyl)-N'-1-piperidinyl- (9CI) (CA INDEX NAME)

Pyrazinecarboxylic acid, 5,6-bis(4-methylphenyl)-3-[(1-

piperidinylamino)carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX

811441-53-9 CAPLUS

811441-54-0 CAPLUS

1-piperidinyl- (9CI) (CA INDEX NAME)

NH- (CH2)3-OH

2,3-Pyrazinedicarboxamide, 5,6-bis(4-chlorophenyl)-N-(3-hydroxybutyl)-N'-1piperidinyl- (9CI) (CA INDEX NAME)

2,3-Pyrazinedicarboxamide, 5,6-bis(4-chlorophenyl)-N-(3-hydroxypropyl)-N'-

811441-52-8 CAPLUS

RN

CN

RN CN

RN

CN

- RN 811441-58-4 CAPLUS
- CN Pyrazinecarboxamide, 5,6-bis(4-methylphenyl)-N-1-piperidinyl-3-(1H-tetrazol-1-ylmethyl)- (9CI) (CA INDEX NAME)

- RN 811441-62-0 CAPLUS
- CN Pyrazinecarboxamide, 5,6-bis(4-methylphenyl)-N-1-piperidinyl-3-(2H-tetrazol-2-ylmethyl)- (9CI) (CA INDEX NAME)

RN 811441-64-2 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-1-piperidinyl-3-(2H-tetrazol-2-ylmethyl)- (9CI) (CA INDEX NAME)

RN 811441-65-3 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-1-piperidinyl-3-(1H-tetrazol-1-ylmethyl)- (9CI) (CA INDEX NAME)

RN 811441-66-4 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-(4,4-difluorocyclohexyl)-3-(2H-tetrazol-2-ylmethyl)- (9CI) (CA INDEX NAME)

RN 811441-67-5 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-(4,4-difluoro-1-piperidinyl)-3-(2H-tetrazol-2-ylmethyl)- (9CI) (CA INDEX NAME)

- 811441-68-6 CAPLUS CN
 - Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-[(2-methoxyethoxy)methyl]-N-1-piperidinyl- (9CI) (CA INDEX NAME)

MeO-CH2-CH2-O-CH2

- RN 811441-71-1 CAPLUS
- Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-[(5-cyclopropyl-2H-tetrazol-2-yl)methyl]-N-1-piperidinyl- (9CI) (CA INDEX NAME) CN

RN 811441-74-4 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-[(5-cyclopropyl-1H-tetrazol-1-yl)methyl]-N-1-piperidinyl- (9CI) (CA INDEX NAME)

RN 811441-75-5 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-[(5-methyl-2H-tetrazol-2-yl)methyl]-N-1-piperidinyl- (9CI) (CA INDEX NAME)

- RN 811441-78-8 CAPLUS
- CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-[(5-methyl-1H-tetrazol-1-yl)methyl]-N-1-piperidinyl- (9CI) (CA INDEX NAME)

- RN 811441-79-9 CAPLUS
- CN Pyrazinecarboxylic acid, 6-(4-chlorophenyl)-5-(4-methylphenyl)-3-[(1-piperidinylamino)carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

- RN 811441-86-8 CAPLUS
- CN Pyrazinecarboxylic acid, 5-(4-chlorophenyl)-6-(4-methylphenyl)-3-[(1-piperidinylamino)carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME)

- RN 811441-87-9 CAPLUS
- CN Pyrazinecarboxamide, 6-(4-chlorophenyl)-5-(4-methylphenyl)-N-1-piperidinyl-3-(2H-tetrazol-2-ylmethyl)- (9CI) (CA INDEX NAME)

- 811441-94-8 CAPLUS RN
- Pyrazinecarboxamide, 5-(4-chloropheny1)-6-(4-methylpheny1)-N-1-piperidinyl-3-(2H-tetrazol-2-ylmethyl)- (9CI) (CA INDEX NAME) CN

$$N = N$$

$$N =$$

- RN 811441-97-1 CAPLUS
- Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[[(2-hydroxyethyl)meino]carbonyl]-, 1,1-dimethylethyl ester (9CI) (CA INDEX NAME) CN

- RN 811441-98-2 CAPLUS
- CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-1-piperidinyl-3-propoxy-(9CI) (CA INDEX NAME)

- RN 811442-03-2 CAPLUS
- CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-1-piperidinyl-3-(1H-tetrazol-5-ylmethyl)- (9CI) (CA INDEX NAME)

RN 811442-07-6 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chloropheny1)-3-[[5-(4-morpholiny1)-2H-tetrazo1-2-y1]methy1]-N-1-piperidiny1- (9CI) (CA INDEX NAME)

RN 811442-08-7 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-[[5-(4-morpholinyl)-1H-tetrazol-1-yl]methyl]-N-1-piperidinyl- (9CI) (CA INDEX NAME)

RN 811442-10-1 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-1-piperidinyl-3-[[5-(1pyrrolidinyl)-2H-tetrazol-2-yl]methyl]- (9CI) (CA INDEX NAME)

RN 811442-11-2 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-1-piperidinyl-3-[[5-(1-pyrrolidinyl)-1H-tetrazol-1-yl]methyl]- (9CI) (CA INDEX NAME)

RN 811442-12-3 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-[[5-(methylthio)-2H-tetrazol-2-yl]methyl]-N-1-piperidinyl- (9CI) (CA INDEX NAME)

811442-13-4 CAPLUS RN CN

Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-[[5-(methylthio)-1H-tetrazol-1-yl]methyl]-N-1-piperidinyl- (9CI) (CA INDEX NAME)

RN 811442-14-5 CAPLUS

Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-(4,4-difluorocyclohexyl)-3-(methoxymethyl)- (9CI) (CA INDEX NAME) CN

RN 811442-16-7 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-(4,4-difluorocyclohexyl)-3-[[(4-fluorophenyl)methoxy]methyl]- (9CI) (CA INDEX NAME)

RN 811442-19-0 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-[(4,4-difluoro-1-piperidinyl)methyl]-N-1-piperidinyl- (9CI) (CA INDEX NAME)

- RN 811442-21-4 CAPLUS
- CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-(4,4-difluorocyclohexyl)-3-[(4,4-difluoro-1-piperidinyl)methyl]- (9CI) (CA INDEX NAME)

- RN 811442-22-5 CAPLUS
- CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-N-(4,4-difluoro-1-piperidinyl)-3-(methoxymethyl)- (9CI) (CA INDEX NAME)

RN 811442-24-7 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chlorophenyl)-3-[[4-(1-hydroxyethyl)-1H-1,2,3-triazol-1-yl]methyl]-N-1-piperidinyl- (9CI) (CA INDEX NAME)

RN 811442-25-8 CAPLUS

CN Pyrazinecarboxamide, 3-[[4-(aminomethyl)-1H-1,2,3-triazol-1-yl]methyl]-5,6-bis(4-chlorophenyl)-N-1-piperidinyl- (9CI) (CA INDEX NAME)

RN 811442-26-9 CAPLUS

CN Pyrazinecarboxamide, 3-[[5-(aminomethyl)-1H-1,2,3-triazol-1-yl]methyl]-5,6-bis(4-chlorophenyl)-N-1-piperidinyl- (9CI) (CA INDEX NAME)

IT 811441-28-8P, Ethyl 3-(azidomethyl)-5,6-bis(4chlorophenyl)pyrazine-2-carboxylate 811441-29-9P, 3-(Azidomethyl)-5,6-bis(4-chlorophenyl)pyrazine-2-carboxylic acid 811441-30-2P, 3-(Azidomethyl)-5,6-bis(4-chlorophenyl)pyrazine-2carboyl chloride 811441-31-3P, 3-(Azidomethyl)-5,6-bis(4chlorophenyl)-N-(piperidin-1-yl)pyrazine-2-carboxamide 811442-09-8P, 5,6-Bis(4-chlorophenyl)-3-(hydroxymethyl)-N-(piperidin-1-yl)pyrazine-2-carboxamide

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(intermediate; preparation of 3-substituted 5,6-diarylpyrazine-2-carboxamide and 2-sulfonamide derivs. as CB1 modulators)

RN 811441-28-8 CAPLUS CN Pyrazinecarboxylic a

N Pyrazinecarboxylic acid, 3-(azidomethyl)-5,6-bis(4-chlorophenyl)-, ethyl

ester (9CI) (CA INDEX NAME)

- RN 811441-29-9 CAPLUS
- CN Pyrazinecarboxylic acid, 3-(azidomethyl)-5,6-bis(4-chlorophenyl)- (9CI) (CA INDEX NAME)

- RN 811441-30-2 CAPLUS
- CN Pyrazinecarbonyl chloride, 3-(azidomethyl)-5,6-bis(4-chlorophenyl)- (9CI) (CA INDEX NAME)

- RN 811441-31-3 CAPLUS
- CN Pyrazinecarboxamide, 3-(azidomethy1)-5,6-bis(4-chloropheny1)-N-1-piperidinyl- (9CI) (CA INDEX NAME)

RN 811442-09-8 CAPLUS

CN Pyrazinecarboxamide, 5,6-bis(4-chloropheny1)-3-(hydroxymethy1)-N-1-piperidiny1- (9CI) (CA INDEX NAME)

(preparation of 3-substituted 5,6-diarylpyrazine-2-carboxamide and 2-sulfonamide derivs. as CB1 modulators)

RN 811441-51-7 CAPLUS

CN Pyrazinecarboxylic acid, 5,6-bis(4-chlorophenyl)-3-[(1-piperidinylamino)carbonyl]- (9CI) (CA INDEX NAME)

REFERENCE COUNT:

6 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT L7 ANSWER 12 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2004:205967 CAPLUS DOCUMENT NUMBER: 142:113926

AUTHOR(S):

TITLE: Product class 14: pyrazines

Sato, N. CORPORATE SOURCE: Germany

SOURCE: Science of Synthesis (2004), 16, 751-844

CODEN: SSCYJ9 PUBLISHER: Georg Thieme Verlag

DOCUMENT TYPE: Journal: General Review LANGUAGE: Enalish

A review. Methods for preparing pyrazines are reviewed including cyclization, ring transformation, aromatization and substituent

modification. тт 64344-98-5P 101445-25-4P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of pyrazines via cyclization, ring transformation, aromatization and substituent modification)

RM 64344-98-5 CAPLUS

CN Pyrazinecarboxamide, 3-(cyclohexylamino)-5,6-diphenyl- (9CI) (CA INDEX NAME)

101445-25-4 CAPLUS RN

Pyrazinecarboxamide, 3-amino-5,6-diphenyl- (9CI) (CA INDEX NAME)

506 THERE ARE 506 CITED REFERENCES AVAILABLE FOR REFERENCE COUNT: THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L7 ANSWER 13 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1997:102094 CAPLUS

DOCUMENT NUMBER: 126:199575

TITLE: Tricyclic substituted hexahydrobenz[e]isoindole

alpha-1 adrenergic antagonists

INVENTOR(S): Meyer, Michael D.; Altenbach, Robert J.; Basha, Fatima Z.; Carroll, William A.; Drizin, Irene; Elmore, Steven W.; Kerwin, James F., Jr.; Lebold, Suzanne A.; Lee, Edmund L.; Sippy, Kevin B.; Tietje, Karin R.; Wendt,

Michael D.

PATENT ASSIGNEE(S): Abbott Laboratories, USA

SOURCE: U.S., 73 pp., Cont.-in-part of U.S. Ser. No. 379,414, abandoned.

CODEN: USXXAM

DOCUMENT TYPE: Patent LANGUAGE: English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.			APPLICATION NO.	DATE
			US 1995-463528	
IL 116405				
			CA 1996-2211212	
			WO 1996-US72	
W: AU, CA,				
RW: AT, BE,	CH, DE, DK	, ES, FR,	GB, GR, IE, IT, LU,	MC, NL, PT, SE
AU 9647457	A	19960814	AU 1996-47457	19960111
AU 705283	B2	19990520		
EP 808318	A1	19971126	EP 1996-903340	19960111
EP 808318	B1	20000628		
R: AT, BE,	CH, DE, DK	, ES, FR,	GB, GR, IT, LI, LU,	NL, SE, PT, IE
AT 194141	T	20000715	AT 1996-903340	19960111
ES 2149451	T3	20001101	ES 1996-903340	19960111
PT 808318	T	20001229	PT 1996-903340	19960111
JP 2001504797	T	20010410	JP 1996-522867	19960111
GR 3034485	T3	20001229	GR 2000-402174	20000926
PRIORITY APPLN. INFO			US 1995-379414	
	-		US 1995-463528	
			WO 1996-US72	
OTHER SOURCE(S):	MARPAT	126:19957		

AB I (W = tricyclic heterocyclic ring system, e. g. pyrazinothienopyrimidinediones, pyridofuropyrimidinediones, pyrazinothienopyrimidinediones; n = 2-6; R1 and R2 = H, alkoxy, hydroxy, alkyl, halo, carboxy, alkoxycarbonyl) and their pharmaceutically acceptable salts were prepared I are $\alpha-1$ adrenergic antagonists and useful in the treatment of BPH (benign prostrate hyperplasia). α -1 Antagonist compns. and a method for antagonizing $\alpha-1$ receptors and treating BPH are also disclosed. 34121-79-4

RL: RCT (Reactant); RACT (Reactant or reagent) (for preparation of tricyclic substituted hexahydrobenzisoindoles as alpha-1 adrenergic antagonists)

RN 34121-79-4 CAPLUS

Ι

CN Pyrazinecarboxamide, 3,4-dihydro-3-oxo-5,6-diphenyl- (9CI) (CA INDEX NAME)

L7 ANSWER 14 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1996:580282 CAPLUS

DOCUMENT NUMBER: 125:221858

TITLE: Preparation of tricyclic substituted benz[e]isoindoles

as αl adrenergic antagonists

INVENTOR(S): Meyer, Michael D.; Altenbach, Robert J.; Basha, Fatima
Z.; Carroll, William A.; Drizin, Irene; Kerwin, James
F., Jr.; Lebold, Suzanne A.; Lee, Edmund L.; Elmore,

Steven W.; et al.

PATENT ASSIGNEE(S): Abbott Laboratories, USA SOURCE: PCT Int. Appl., 180 pp.

CODEN: PIXXD2
DOCUMENT TYPE: Patent
LANGUAGE: English

FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO.	DATE
WO 9622992 W: AU, CA, JP,		0801 WO 1996-US72	19960111
		FR, GB, GR, IE, IT, LU,	MC, NL, PT, SE
US 5597823	A 1997	0128 US 1995-463528	19950605
AU 9647457	A 1996	0814 AU 1996-47457	19960111
AU 705283	B2 1999	0520	
EP 808318	A1 1997	1126 EP 1996-903340	19960111
EP 808318	B1 2000	0628	
R: AT, BE, CH,	DE, DK, ES,	FR, GB, GR, IT, LI, LU,	NL, SE, PT, IE
AT 194141		0715 AT 1996-903340	19960111
JP 2001504797	T 2001	0410 JP 1996-522867	19960111
GR 3034485	T3 2000	1229 GR 2000-402174	20000926
PRIORITY APPLN. INFO.:		US 1995-379414	A 19950127
		US 1995-463528	A 19950605
		WO 1996-US72	W 19960111
OTUED COMPCE/C).	MADDAT 125.	221050	

OTHER SOURCE(S): MARPAT 125:221858

- * STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY AVAILABLE VIA OFFLINE PRINT *
- AB The title compds. [I; Rl, R2 = H, alkoxy, OH, etc.; W = tricyclic heterocyclic ring system; n = 2-6] and their salts, useful in the treatment of benign prostatic hypertrophy (BPH), were prepared Thus, reaction of urea II with benz[e]isoindole III in the presence of (Pr)2NEt in DMSO afforded the desired product cis-IV.RCl which showed pA2 of 8.37 for inhibition of phenylepherine(PE)-induced contraction of rat vas.

 II 34121-79-4
 - RI: RCT (Reactant); RACT (Reactant or reagent) (preparation of tricyclic substituted benz[e]isoindoles as αl adrenerqic antaqonists)

CN Pyrazinecarboxamide, 3,4-dihydro-3-oxo-5,6-diphenyl- (9CI) (CA INDEX NAME)

L7 ANSWER 15 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1996:466654 CAPLUS

DOCUMENT NUMBER: 125:157774

TITLE: Anthelmintic activity of 6,7-diarylpteridines

AUTHOR(S): Ochoa, Carmen; Rodriguez, Juan; Lopez Garcia, Maria Luz; Martinez, Antonio Ramon; Martinez, Maria Mercedes

CORPORATE SOURCE: Fac. Farm., Univ. Complutense, Madrid, E-28006, Spain SOURCE: Arzneimittel-Forschung (1996), 46(6), 643-648

CODEN: ARZNAD: ISSN: 0004-4172

PUBLISHER: Cantor

DOCUMENT TYPE: Journal LANGUAGE: English

AB In search for new anthelmintic compds., some 6,7-diaryl-pteridines were synthesized from the corresponding diaminopyrimidines and aromatic aldehydes. Their anthelmintic activity was tested in vitro against Caenorhabditis elegans and Heligmosomoides polygyrus and in vivo against Trichinella spiralis. Structure-activity relationships are discussed.

IT 180603-98-9P 180603-99-0P

RI: BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); BIOL (Biological study); PREP (Preparation)

(anthelmintic activity and preparation of diarylpteridines)

RN 180603-98-9 CAPLUS

CN Pyrazinecarboxamide, 3-amino-5,6-di-2-thienyl- (9CI) (CA INDEX NAME)

RN 180603-99-0 CAPLUS

CN Pyrazinecarboxamide, 3-amino-N-butyl-5,6-di-2-thienyl- (9CI) (CA INDEX NAME)

L7 ANSWER 16 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1987:119534 CAPLUS

DOCUMENT NUMBER: 106:119534

TITLE: Pteridines. LXXVIII. Reactions and properties of 4-thiolumazine derivatives

AUTHOR(S): Lutz, Herman; Pfleiderer, Wolfgang

CORPORATE SOURCE: Fak. Chem., Univ. Konstanz, Konstanz, D-7750, Fed. Rep. Ger.

SOURCE: Croatica Chemica Acta (1986), 59(1), 199-220

CODEN: CCACAA; ISSN: 0011-1643

English

DOCUMENT TYPE:

LANGUAGE:

AB The 4-thioxo function in the 6,7-diphenyl-4-thiolumazines I (X = S, R, R1 = H, Me) showed easy displacement by nucleophiles under mild conditions. Special structural and electronic features became obvious with I (X = S, R = H, R1 = Me), which reacted analogously to I (X = S, R = R1 = Me) with amines to I (X = NH, NNe, NET, NBu, NNHPh, NHMMeh). The latter compds. are very light-sensitive and react by photooxidn. to give I (X = O). Nucleophilic displacement by alkoxides under HgBr2 catalysis yielded the unusual 4,4-di-O-alkyl acetals I [X = (OMe)2, OCH2CH2O]. The acetal function is prone to easy substitution by C-H acidic compds., giving I [X = C(CR)2] from I [X = (OMe)2).

T 25472-83-7P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 25472-83-7 CAPLUS

CN Pyrazinecarboxamide, N-methyl-3-(methylamino)-5,6-diphenyl- (8CI, 9CI) (CA INDEX NAME)

L7 ANSWER 17 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1982:68939 CAPLUS DOCUMENT NUMBER: 96:68939

ORIGINAL REFERENCE NO.: 96:11329a,11332a

TITLE: Synthesis of pyrazinedicarboximides from

diaminomaleonitrile
AUTHOR(S): Tsuda, Tadataka; Fujishima, Katsuhiro; U

AUTHOR(S): Tsuda, Tadataka; Fujishima, Katsuhiro; Ueda, Hiroo CORPORATE SOURCE: Coll. Agric., Univ. Osaka Prefect., Osaka, 591, Japan SOURCE: Agricultural and Biological Chemistry (1981), 45(9),

> 2129-30 CODEN: ABCHA6; ISSN: 0002-1369

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 96:68939

GT

- AB Hydrolysis of pyrazines I (R = H, Me, Ph, 4-ClC6H4, 3,4-Cl2C6H3, 4-MeC6GH4; RI = H, Me, Ph, R2 = CN), prepared from diaminomaleonitrile, followed by esterification gave I (R2 = CO2Me)(II). Amidn. of II with NH3 followed by intramol. cyclocondensation gave the title compds. (III). II (R = Ph, R1 = H, R2 = CO2Me) showed bactericidal activity superior to that of phenazine oxide.
- IT 80356-91-8P
 RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT
 (Reactant or reagent)
 (preparation and cyclization of, pyridinedicarboximide from)
- RN 80356-91-8 CAPLUS
 CN 2,3-Pyrazinedicarboxamide, 5,6-diphenyl- (CA INDEX NAME)

L7 ANSWER 18 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1980:586294 CAPLUS

DOCUMENT NUMBER: 93:186294

ORIGINAL REFERENCE NO.: 93:29698h,29699a

TITLE: One-step preparation of 3-alkoxypyrazine-2-

carbonitriles from pyrazine-2,3-dicarbonitriles and

related reactions

AUTHOR(S): Kojima, Takakazu; Nagasaki, Fumihiko; Ohtsuka, Yozo CORPORATE SOURCE: Fine Chem. Res. Lab., Nippon Soda Co. Ltd., Odawara,

250-02, Japan

SOURCE: Journal of Heterocyclic Chemistry (1980), 17(3), 455-9

CODEN: JHTCAD; ISSN: 0022-152X

DOCUMENT TYPE: Journal

LANGUAGE: English

OTHER SOURCE(S): CASREACT 93:186294

GT

AB Disubstituted alkoxypyrazinecarbonitriles I (R = Ph, H, 1,8-C10H6,

9,10-phenanthrenediyl; R1 = alkyl) were prepared from the pyrazinedicarbonitriles II by direct substitution with alcs. Treatment of II with amines gave either pyrrolopyrazines III (R = H, Ph) or

substitution products. In a low temperature range, II afforded imidates and related compds. The preference among these reactions depended on the 5,6-substituents and on the reaction conditions.

IT 75018-16-5P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 75018-16-5 CAPLUS
CN Pyrazinecarboximida

Pyrazinecarboximidamide, N-butyl-3-(butylamino)-5,6-diphenyl- (9CI) (CA INDEX NAME)

Ph N C-NHBu-n

L7 ANSWER 19 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1980:446712 CAPLUS

ACCESSION NUMBER: 1980:446712 CAPLU DOCUMENT NUMBER: 93:46712

ORIGINAL REFERENCE NO.: 93:7730h,7731a

TITLE: Pyrazinecvanocarboxamides

INVENTOR(S): Genda, Yoshikazu; Tomita, Nobuo; Ito, Masaru; Kano,

PATENT ASSIGNEE(S): Nippon Soda Co., Ltd., Japan SOURCE: Jpn. Kokai Tokkyo Koho, 3 pp.

CODEN: JKXXAF DOCUMENT TYPE: Patent Japanese

LANGUAGE: FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
JP 54154776	A	19791206	JP 1978-63655	19780527
JP 61056230	В	19861201		
PRIORITY APPLN. INFO.:			JP 1978-63655 A	19780527

Title compds. I (R = H, Me, Ph) were prepared by treating II with HCl and AB AcOH. Thus, stirring a mixture of 5 g II, 40 mL 35% HCl, and 5 mL AcOH for 3 h 15 min at 30-5° gave 86.1% I (R = H).

66371-68-4P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

66371-68-4 CAPLUS

RN

CN Pyrazinecarboxamide, 3-cyano-5,6-diphenyl- (9CI) (CA INDEX NAME)

L7 ANSWER 20 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1980:41887 CAPLUS

DOCUMENT NUMBER: 92:41887 ORIGINAL REFERENCE NO.: 92:6993a,6996a

TITLE: Chemistry of diaminomaleonitrile. 5. Dihydropyrazine synthesis

AUTHOR(S): Ohtsuka, Yozo; Tohma, Eiko; Kojima, Sigeru; Tomita,

CORPORATE SOURCE: Sagami Chem. Res. Cent., Sagamihara, 229, Japan

SOURCE: Journal of Organic Chemistry (1979), 44(26), 4871-6 CODEN: JOCEAH; ISSN: 0022-3263

DOCUMENT TYPE: Journal

LANGUAGE: English

CASREACT 92:41887 OTHER SOURCE(S):

- AB Condensation of RCHO (R = optionally substituted Ph) with Schiff bases I (R1 = optionally substituted Ph, CHMe2) in the presence of NEt3 <20° is accompanied by regiospecific hydration of the nitrile groups to give 3-cyanoacrylamide derivs. II, which cyclize readily into 1,2-dihydropyrazines III and IV. The substituent effect on the product ratio is examined, and the reaction mechanism is discussed in terms of a new general reaction pattern of diaminomaleonitrile derivative Reactions of III and IV by oxidation, reduction, hydantoin formation with isocyanates, and cyanoethylation are also reported.
 - I 66371-68-4P 71871-19-7P 71871-20-0P 71871-22-2P 71871-23-3P 71871-24-4P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)
- RN 66371-68-4 CAPLUS
- CN Pyrazinecarboxamide, 3-cyano-5,6-diphenyl- (9CI) (CA INDEX NAME)

- RN 71871-19-7 CAPLUS
- CN Pyrazinecarboxamide, 3-cyano-6-(4-methylphenyl)-5-phenyl- (9CI) (CA INDEX NAME)

- RN 71871-20-0 CAPLUS
- CN Pyrazinecarboxamide, 3-cyano-5-(4-methylphenyl)-6-phenyl- (9CI) (CA INDEX NAME)

71871-22-2 CAPLUS RN

CN Pyrazinecarboxamide, 3-cyano-6-(4-nitrophenyl)-5-phenyl- (9CI) (CA INDEX NAME)

RN 71871-23-3 CAPLUS

CN Pyrazinecarboxamide, 3-cyano-6-(4-cyanophenyl)-5-phenyl- (9CI) (CA INDEX NAME)

RN 71871-24-4 CAPLUS

CN Pyrazinecarboxamide, 3-cyano-6-(4-cyanophenyl)-5-(4-methylphenyl)- (9CI) (CA INDEX NAME)

ANSWER 21 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1978:171793 CAPLUS

DOCUMENT NUMBER: 88:171793

ORIGINAL REFERENCE NO.: 88:27075a,27078a TITLE:

1,2-Dihydropyrazine derivatives INVENTOR(S):

Ohtsuka, Yozo; Ito, Masaru; Tomita, Nobuo

PATENT ASSIGNEE(S): Nippon Soda Co., Ltd., Japan; Sagami Chemical Research

Center SOURCE: Ger. Offen., 48 pp.

CODEN: GWXXBX

DOCUMENT TYPE: Patent LANGUAGE: German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE
DE 2736230	A1	19780216	DE 1977-2736230	19770811
JP 53022529	A	19780302	JP 1976-96020	19760813
JP 57045260	В	19820927		
PRIORITY APPLN. INFO.:			JP 1976-96020 #	19760813
CT				

AB Title compds. (I; R, R1 = Ph, condensed aromatic, or heterocyclic groups), fast yellow dyes showing a green to yellow luminescence, are prepared (a) by condensing RCH:NC(CN):C(CN)NH2 with R1CHO in the presence of base to give RCH:NC(CN):C(CONH2)N:CHR1, followed by ring closure, or (b) by selective hydrolysis of II to III, followed by selective reduction Thus, reaction of PhCH:NC(CN):C(CN)NH2 [56029-18-6] with PhCHO [100-52-7] in EtOH containing Et3N gave PhCH:NC(CN):C(CONH2)N:CHPh [66371-72-0], which was cyclized by heating with Me2SO to form a mixture of IV [66371-73-1] and V [66371-74-2]. The IV-V mixture, resolvable by fractional recrystn., showed (Japanese standard test K 5101) a brilliant greenish yellow tone, solvent stability 4-5 (1 lowest, 5 highest), and water stability 5, and lightfastness (Fade-O-meter) 7-8.

66371-68-4P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and selective reduction of)

RN 66371-68-4 CAPLUS

Pyrazinecarboxamide, 3-cyano-5,6-diphenyl- (9CI) (CA INDEX NAME) CN

L7 ANSWER 22 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1977:552132 CAPLUS

DOCUMENT NUMBER: 87:152132

ORIGINAL REFERENCE NO.: 87:24075a,24078a

TITLE: Amidinoacetamides in the synthesis of pyrazines and pteridines

AUTHOR(S): Keir, William F.; MacLennan, Alexander H.; Wood,

Hamish C. S.

CORPORATE SOURCE: Paislev Coll. Technol., Paislev, UK

SOURCE: Journal of the Chemical Society, Perkin Transactions

1: Organic and Bio-Organic Chemistry (1972-1999)

(1977), (11), 1321-5

CODEN: JCPRB4; ISSN: 0300-922X

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 87:152132

GI

O N. Me

AB Cyclocondensation of 2-(substituted amidino)-2-aminoacetamides with 1,2-dicarbonyl compds. gave 3-(substituted amino)-pyrazine-2-carboxamides which with one-carbon units gave 1-substituted pteridin-4(1R-)-ones and -2,4-(1R)-diones. E.g., PhcE2NHC(:NH)CH(NH2)CONH2.HCl with biacetyl gave 80% pyrazine I which with HCO2H and ClCO2Et gave 60% pteridinone III and 59% pteridinedione III, resp.

IT 64344-98-5P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and cyclocondensation reaction of, with

diethoxydimethylformamide)

RN 64344-98-5 CAPLUS

CN Pyrazinecarboxamide, 3-(cyclohexylamino)-5,6-diphenyl- (9CI) (CA INDEX NAME)

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and cyclocondensation reactions of, with formic acid and Et chloroformate)

64344-96-3 CAPLUS

N Pyrazinecarboxamide, 5,6-diphenyl-3-[(phenylmethyl)amino]- (9CI) (CA INDEX NAME)

L7 ANSWER 23 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1976:593017 CAPLUS

DOCUMENT NUMBER: 85:193017 ORIGINAL REFERENCE NO.: 85:30879a

ORIGINAL REFERENCE NO.: 85:30879a,30882a
TITLE: Nucleosides, XIX.

TITLE: Nucleosides, XIX. Synthesis, properties and chemical behavior of 1(3)-methyl-6,7-diphenyl-3(1)-(β -D-

ribofuranosyl)lumazine derivatives

AUTHOR(S): Kobayashi, Kiyotaka; Pfleiderer, Wolfgang

CORPORATE SOURCE: Fachber. Chem., Univ. Konstanz, Konstanz, Fed. Rep.

SOURCE: Chemische Berichte (1976), 109(9), 3194-207

CODEN: CHBEAM: ISSN: 0009-2940

DOCUMENT TYPE: Journal LANGUAGE: German

GI For diagram(s), see printed CA Issue.

AB Ribofuranosyllumazine I (R = R2, R1 = Me, R3-R5 = H) (II) was prepared by coupling 1-O-acetyl-2,3,5-tri-O-benzoyl-β-D-ribofuranose (III) with

O-trimethylsilyl derivative of I (R = H, R1 = Me) followed by alkaline hydrolysis.

Similarly I (R = Me, Rl = R2, R3-R5 = H) (IV) was prepared from I (R = Me, Rl = H) and III. Isopropylidenation of II and IV gave I (R = R2, Rl = H, R4R5 = CMe2) (V) and I (R = H, Rl = R2, R4R5 = CMe2) (VI). In the alkaline hydrolysis of IV-VI the nucleophilic attack occurred at the CO group at C-2 with cleavage of the pyrimidine ring and formation of the corresponding 3-amino-5,6-diphenyl-2-pyrazinecarboxamides.

IT 25472-83-7P

RL: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation); RACT (Reactant or reagent)

(preparation and reaction of, with ethyl chloroformate)

RN 25472-83-7 CAPLUS

Pyrazinecarboxamide, N-methyl-3-(methylamino)-5,6-diphenyl- (8CI, 9CI) (CA INDEX NAME)

60980-87-2P 60980-97-4P 60980-98-5P 60980-99-6P 60981-00-2P 60981-01-3P 60981-02-4P 60981-03-5P 60981-04-6P 60981-05-7P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 60980-87-2 CAPLUS

Pyrazinecarboxamide, N-methyl-5,6-diphenyl-3-[(2,3,4-tri-0-acetyl- α -D-ribopyranosyl)aminol- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

RN 60980-97-4 CAPLUS

CN Pyrazinecarboxamide, N-methyl-5,6-diphenyl-3-(D-ribofuranosylamino)- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

60980-98-5 CAPLUS

CN Pyrazinecarboxamide, 3-amino-N-methyl-5,6-diphenyl- (9CI) (CA INDEX NAME)

RN 60980-99-6 CAPLUS

CN Pyrazinecarboxamide, N-methyl-5,6-diphenyl-3-(α-Dribopyranosylamino) - (9CI) (CA INDEX NAME)

Absolute stereochemistry.

- RN 60981-00-2 CAPLUS
- CN Pyrazinecarboxamide, N-methyl-5,6-diphenyl-3-[(2,3,4-tri-0-acetyl-β-D-ribopyranosyl)amino]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

- RN 60981-01-3 CAPLUS
- CN Pyrazinecarboxamide, N-methyl-5,6-diphenyl-3-[(2,3,5-tri-0-acetyl- α -D-ribofuranosyl)amino]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

- RN 60981-02-4 CAPLUS
- CN Pyrazinecarboxamide, N-methyl-5,6-diphenyl-3-[(2,3,5-tri-O-acetyl-β-D-ribofuranosyl)amino]- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

- RN 60981-03-5 CAPLUS
- CN Pyrazinecarboxamide, N-methyl-3-[[2,3-0-(1-methylethylidene)-β-D-ribofuranosyl]amino]-5,6-diphenyl- (9CI) (CA INDEX NAME)

Absolute stereochemistry.

- RN 60981-04-6 CAPLUS
- CN Carbamic acid, methyl[3-[[[2,3-0-(1-methylethylidene)-β-D-ribofuranosyl]amino|carbonyl]-5,6-diphenylpyrazinyl]-, monosodium salt (9CI) (CA INDEX NAME)

Absolute stereochemistry.

Na

- RN 60981-05-7 CAPLUS
- CN Carbamic acid, methyl[3-[(methylamino)carbonyl]-5,6-diphenylpyrazinyl]-, ethyl ester (9CI) (CA INDEX NAME)

L7 ANSWER 24 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1971:518287 CAPLUS

DOCUMENT NUMBER: 75:118287 ORIGINAL REFERENCE NO.: 75:18673a,18676a

TITLE:

Alkylation of 4-oxopteridines

AUTHOR(S): CORPORATE SOURCE: SOURCE:

DOCUMENT TYPE:

Neiman, Zohar; Bergmann, Felix; Meyer, Amatzya Y. Hadassah Med. Sch., Hebrew Univ., Jerusalem, Israel

Chem. Biol. Pteridines, Proc. Int. Symp., 4th (1970), Meeting Date 1969, 29-34. Editor(s): Iwai, K. Int.

Acad. Print. Co.: Tokvo, Japan.

CODEN: 23BVAJ Conference English

LANGUAGE: For diagram(s), see printed CA Issue.

4-Pteridin-one (I), 6,7-dimethyl-4-pteridinone (II), and

6,7-diphenyl-4-pteridinone (III) were alkylated exclusively in the pyrimidine ring by MeI-DMF to yield the corresponding 1,3-dimethyl-4oxopteridinium salts IV, V, and VI in 10%, 50% and 50% yield, resp. The pyrimidine ring of these methylation products was cleaved readily by hot 2N NaOH to yield the corresponding pyrazines. Reduction of IV, V, and VI with NaBH4 yielded the corresponding derivs. of 1,2-dihydropteridine. The reaction path to IV, V, and VI was studied by paper chromatog., and

related with charge ds. calculated by the HMO and the SCF-Pariser-Pople-Parr

methods

25472-83-7P RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) RN 25472-83-7 CAPLUS

CN Pyrazinecarboxamide, N-methyl-3-(methylamino)-5,6-diphenyl- (8CI, 9CI) (CA INDEX NAME)

ANSWER 25 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1971:488570 CAPLUS DOCUMENT NUMBER: 75:88570

ORIGINAL REFERENCE NO.: 75:14029a,14032a

TITLE: New oral antidiabetic drugs. I

Ambrogi, V.; Bloch, Konrad; Daturi, S.; Griggi, P.; AUTHOR(S): Logemann, W.; Parenti, M. A.; Rabini, T.; Tommasini, R.

CORPORATE SOURCE: Ist. Carlo Erba Ric. Ter., Milan, Italy SOURCE: Arzneimittel-Forschung (1971), 21(2), 200-4

CODEN: ARZNAD; ISSN: 0004-4172
DOCUMENT TYPE: Journal

LANGUAGE: English

GI For diagram(s), see printed CA Issue.

AB All of 20 new pyrazinecarboxamidoethylphenylnesulfonylureas had

hypoglycemic activity in mice, and 19 were active in rats; in rats N - (4 - [β - (5 -methylpyrazine -2-carboxamido)ethyl]phenylsulfonyl)-N'-

cyclohexylurea (I) was the most active producing a hypoglycemic activity of 46% at 1.5 $\,$ mg/kg orally. 4-(4-[β -(5-Methylpyrazine-2-

carboxamido)ethyllphenylsulfonyl)-1,1 - hexamethylenesemicarbazide (II), the only pyrazinecarboxamidoethylphenylsulfonylsemicarbazide tested, was as effective as I at the same dose. Neither of the 2

as effective as 1 at the same dose. Neither of the 2 pyrazinecarboxamidocycloalkylphenylsulfonylureas tested had antidiabetic activity in mice or rats. The sulfonamide were synthesized by reacting pyrazine-, pyridazine-, or pyrimidinecarboxamidobenzenesulfonamides with cyclohexyl isocyanate. Intermediate benzenesulfonamides were prepared by acylation of p-(β -aminoethyl)benzenesulfonamide. II was prepared from Me-4-(β -(β -methylpyrazine-2-carboxamido)ethyl)

phenylsulfonylcarbamate and 1-aminohexamethyleneimine.

IT 33282-78-9P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

RN 33282-78-9 CAPLUS

CN Urea, 1-[[p-[2-(3-amino-5,6-diphenylpyrazinecarboxamido)ethyl]phenyl]sulfo nyl]-3-cyclohexyl- (8CI) (CA INDEX NAME)

L7 ANSWER 26 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1970:55408 CAPLUS DOCUMENT NUMBER: 72:55408

ORIGINAL REFERENCE NO.: 72:10145a,10148a
TITLE: Reduction of quaternary pteridines and purines by

sodium borohydride

AUTHOR(S): Neiman, Zohar

CORPORATE SOURCE: Hadassah Med. Sch., Hebrew Univ., Jerusalem, Israel
SOURCE: Journal of the Chemical Society [Section] C: Ordanic

(1970), (1), 91-4

CODEN: JSOOAX; ISSN: 0022-4952

DOCUMENT TYPE: Journal LANGUAGE: English

OTHER SOURCE(S): CASREACT 72:55408

B In the 3,4-dihydro-1,3-dimethyl-5,6-diphenyl-4-oxopteridinium cation, and in the 1,3-dimethyl-8-phenylhypo-xanthinium cation, position 2 of the pyr imidine ring is reduced by NaBH4. The analogous reaction at position 8 was observed for the 7,9-dimethylhypoxanthinium cation. The structures assigned to the reduction products are supported by spectral data and by

degradation reactions. 25472-83-7P

RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of) RN 25472-83-7 CAPLUS

L7 ANSWER 27 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1968:436172 CAPLUS

DOCUMENT NUMBER: 69:36172

ORIGINAL REFERENCE NO.: 69:6762h,6763a

TITLE: (3-Amino-2-pyrazinecarbonyl)guanidines

SOURCE: U.S., 26 pp.
CODEN: USXXAM
DOCUMENT TYPE: Patent

LANGUAGE: English FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE

US 3313813
DE 1795438
DE DE

APPLICATION NO. DATE

19670411
US 1963-313315
DE

GI For diagram(s), see printed CA Issue.
AB Title compds. I are prepared from II, III, and IV. Thus, 3318 g. SO2C12 is added in 30 min. to 765 g. Me 3-amino-2-pyrazinecarboxylate in 5.1 CGHG, the mixture is agitated 1 hr., refluxed 5 hrs., and agitated overnight to

give 724 g. Me 3-amino-5,6-dichloropyrazinecarboxylate (V), m. 233-4° (MeCN). A mixture of 100 g. V. and 1.1 Me2SO is heated to

 65° and NH3 gas is introduced into the mixture in 45 min. at $65-70^{\circ}$; the mixture is cooled to 10° and NH3 is introduced in

1.25 hrs. to give 91.5% Me 3,5-diamino-6-chloropyrazinecarboxylate, m.

212-13° (MeCN). Also prepared, by known methods are the following II (X, Y, Z, and m.p. given): MeO, NH2, H, 252-4° (decomposition); MeO,

NH2, Br, 217-19°; MeO, NH2, iodine, 200-2°; MeO, PhNH, Cl,

171.5-73°; MeO, p-ClC6H4NH, Cl, 207-8°; MeO, Me2N, Cl, 145.5-6.5°; MeO, MeS, Cl, 214-16°; MeO, MeSO, Cl,

237.5-40.5° (decomposition); MeO, OH, Cl, .apprx.245° (decomposition);

MeO, OH, H, 220-60° (decomposition); MeO, NH2, H, 252-4° (decomposition); MeO, MeO, H, 242.5-3.5°; MeO, MeO, H,

205.5-7.5°; MeO, PhCH2NH, H, 157-8°; MeO, MeO, MeO, Cl, 255-7°; MeO, MeS, Cl, 212-14°; MeO, SH, Cl, 207-8°

(decomposition); MeO, EtO, Cl, 123-5°; MeO, H, Me, 138.5-40.5°; MeO, Cl, Me, 176.5-9.5°; MeO, Me2N, Me, 108.5-10.5°; MeO,

Me, H, 165-7°; MeO, Me, Br, 179-81°; NH2, H, Et,

165.5-8.5°; OH, H, Et, 149-52°; MeO, H, Et, 85-7.5°;

OH, cyclohexyl, H, 182.5-3.5°; MeO, cyclohexyl, H, 173-4.5°; NH2, H, cyclohexyl, -; OH, H, cyclohexyl, -; MeO, H, cyclohexyl,

126.5-8.0°; NH2, H, cyclopropyl, 185.5-7.5°; OH, H, cyclopropyl, 169-72°; MeO, H, cyclohexyl, 112.5-14.5°; MeO,

Ph, H, 231-2°; MeO, H, Ph, 140-1°; MeO, Cl, Ph,

187.5-91.5°; MeO, Ph, Br, 217-21°; OH, H, p-C1C6H4, 213-15°; MeO, H, p-C1C6H4, 181.5-3.5°; MeO, C1, Ph,

213-15°; MeO, H, p-C1C6H4, 181.5-3.5°; MeO, C1, Ph

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187.5-90.5°; MeO, Me2N, Ph, 167-9.5°; MeO, H, Cl,
142° (decomposition); MeO, MeHN, C1, 221-2°; MeO, EtNH, C1,
149-50°; MeO, PrNH, Cl, 138-40°; MeO, iso-PrNH, Cl,
125.5-6.5°; MeO, CH2:CHCH2NH, C1, 105-6.5°; MeO, BuNH, C1,
140-2°; MeO, sec-BuNH, C1, 106-8°; MeO, iso-BuNH, C1,
113.5-15.5°; MeO, tert-BuNH, Cl, 98-108°; MeO, Me(CH2)4NH,
Cl, 100.5-2.5°; MeO, BuCHMeNH, Cl, -; MeO, Et2CHNH, Cl, -; MeO,
Me(CH2)5NH, Cl, 72.5-5.5°; MeO, cyclopropylmethylamino, Cl,
132-3°; MeO, cyclopropylamino, Cl, 167-9°; MeO,
cyclopentylamino, Cl, 119.5-21.5°; MeO, PhCH2NH, Cl, 157-8°;
MeO, p-MeC6H4CH2NH, Cl, 112.5-14.5°; MeO, o-FC6H4CHNH, Cl,
171-4°; MeO, p-C1C6H4CH2NH, C1, 136-7°; MeO, PhCH2CH2NH, C1,
115-19°; MeO, F3CCH2NH, Cl, 153-4°; MeO, F3CCH2CH2NH, Cl,
124.5-5.5°; MeO, HOCH2CH2NH, Cl, 155-7°; MeO,
HOCH2(CHOH) 4CH2NH, Cl, 172-5°; MeO, H2NCH2CH2NH, Cl, 265°;
MeO, Me2NCH2CH2NH, Cl, 257°; MeO, 4-pyridylmethylamino, Cl,
95-7°; Me, 2-furylmethylamino, Cl, 148-9°; MeO, MeEtN, Cl,
102-4°; MeO, MePrN, Cl, 83.5-5.5°; MeO, iso-PrMeN, Cl,
75.5-7.5°; MeO, Me(CH2:CHCH2)N, Cl, 90.5-2°; MeO, MeBun, Cl,
59.5-61.5°; MeO, Et2N, Cl, 99-101°; MeO, EtPrN, Cl, -; MeO,
iso-PrEtN, Cl, -; MeO, Et(CH2:CHCH2)N, Cl, -; MeO, EtBun, Cl,
77.5-9.5°; Me, Pr2N, Cl, 68.5-71.5°; MeO, PrBuN, Cl, -; MeO,
1-pyrrolidinyl, Cl, 168-71°; MeO, hexamethylenimino, Cl,
109-11°; MeO, 4-methylpiperazino, Cl, 186-8°; MeO, MeNHNH,
Cl, 136.5-8°; MeO, Me2NCH2CH2O, Cl, 134.5-6.5°; NH2, H, Cl,
227-30°; OH, H, MeSO2, 239-42° (decomposition).
p-Methylbenzylamine is treated with H2NC(:NH)SMe.0.5H2SO4 to give 28%
p-MeC6H4CH2NHC(:NH)NH2HCl, m. 153-5°. Similarly prepared are
Me(PhCH2)NC(:NH)NH2.HC1, m. 122.5-5.5°, and the following
RNHC(:NH)NH2.HCl (R and m.p. given): o-ClC6H4CH2, 131-6°;
p-C1C6H4CH2, 162.5-4.5°; p-MeOC6H4CH2, 132-7°;
2,4-Me2C6H3CH2, 105-15°; 2,4-C12C6H3CH2, 145-8°;
3,4-Cl2C6H3CH2, 153-7°; PhCH2CH2, 135-8°; PhCH2,
175-8°. 5,6-Diaminouracil-HCl (17.9 g.) is treated at 60°
with 14.9 g. cyclohexylglyoxal-0.5H2O to give 7.5 g. 7-cyclohexyllumazine
[III (X = H, Y = cyclohexyl)], m. 229-31°, which is hydrolyzed to
qive II (X = OH, Y = cyclohexyl, Z = H). Similarly prepared are (m.p.
given): III (X = Me, Y = Ph) [or III (X = Me, Y = Me)], 281.5-2.5°;
III (X = Ph, Y = Me) [or III (X = Me, Y = Ph) [sic], 254.5-5.5°; II
(X = OH, Y = Ph, Z = Me) [or II (X = OH, Y = Me, Z = Ph)],
193.5-4.5°; II (X = OH, Y = Me, Z = Ph) [or II (X = OH, Y = Ph, Z =
Me)] [sic], 155-6^{\circ}. II (X = MeO, Y = Ph, Z = Me) [or II (X = MeO,
Y = Me, Z = Ph) (m. 163-4°) and II (X = MeO, Y = Me, Z = Ph) (or
II (X = MeO, Y = Ph, Z = Me) [sic] (m. 162.5-3.5°) are prepared by
esterification. Methyl 3-isopropylidenamino-6-anilino-2-
pyrazinecarboxylate, m. 195.5-7.50, is prepared from Me2CO and the
amine. Me 3-amino-5,6,7,8-tetrahydroquinoxaline-2-carboxylate, m.
154-5°, and Me 3-amino-7-chloroquinoxaline-2-carboxylate, m.
224.5-5.5°, are prepared by esterification. Alloxan-H2O (61.44 g.)
is treated with 60 q. 3,4-(H2N)2C6H3C1 to give 33% 8-chloroalloxazine, m.
365-6°, and 42% 7-Chloroalloxazine, m. >380°, which is
treated at 165° with NH3 in an autoclave to give 68%
3-amino-7-chloroquinoxaline-2-carboxylic acid, m. 191-2°
(decomposition). A mixture of 33 g. II (X = NH2, Y = H, Z = Cl), 200 ml. Ac20,
and 200 ml. HC(OEt)3 is refluxed 1.5 hrs. to give 20 g.
4-hydroxy-6-chloropteridine (VI), m. 268-70° (decomposition). VI (5.5
g.) is treated with 4.4 g. PhCH2SH to give 5.5 g. 4-hydroxy-6-
benzylthiopteridine (VIII), m. 233-5°. Similarly prepared is
4-hydroxy-6-methylthiopteridine, m. 289.5-91.5°. VII is heated
with NaOH to give II (X = OH, Y = H, Z = PhCH2S(VIII), m. 138.9°.
Similarly prepared is II (X = OH, Y = H, Z = MeS), m. 182-4^{\circ}
(decomposition). II (X = MeO, Y = Me2N, Z = C1) (11.5 g.) is treated with 26.3
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q. H2NC(:NH)NH2.HCl (IX) in the presence of 5.75 q. Na to give 93%
(3-amino-5-dimethylamino-6-chloro-2-pyrazinecarbonyl)quanidine (X), m.
216-17°, HCl salt m. 298° (decomposition). Similarly prepared is
I.HCl (R = R1 = H, X = Y = Cl) (m. 259-61°) which is treated with
Me2NH to give X. II (X = MeO, Y = Me2NCH2CHO, Z = C1) (9.4 g.) is treated
with 20.0 q. IX in the presence of 4 q. Na to give 2.5 q. I.2HCl [R = R1 =
H, X = NHC(:NH)NH2, Z = Cl], m. >340°. A solution of 8.5 5. VIII in
50 ml. Ac20 is heated 5 hrs. to give 6.6 g. 2-methyl-6-benzylthio-4H-
pvrazine[2,3-d][1,3]oxazin-4-one[IV (X = PhCH2S)] (XI), m.
116.5-18.5°; similarly prepared is IV (X = MeS), m. 189-91°.
XI (3.4 g.) is treated with 5.0 g. IX in the presence of 1.0 g. Na to give
1.1 q. I (R = R1 = X = H, Y = PhCH2S), m. 171-3^{\circ} (decomposition). Also
prepared, by the above or related methods, are the following I (R = R1 = H)
(X, Y, and m.p. given): NH2, Br, 232.5-5.5° (decomposition); NH2,
iodine, 273-4° (decomposition); H, MeS, 203-5°; H, MeSO2,
224-6° (decomposition); OH, H, >310°; NH2, H, 286-8°;
Me2N, H, 224-5°; MeO, H, 229-30°; PhCH2NH, H, 231-3°;
the following I (R = R1 = H, Y, = C1) (X and m.p. given): NH2,
240.5-1.5° (HCl salt m. 293.5°); MeNH, 238-9°; EtNH,
217-18°; PrNH, 221-2°; iso-PrNH, 215°; CH2:CHCH2NH,
213-14°; BuNH, 219.5°; sec-BuNH, 208-9°; iso-BuNH,
221°; tert-BuNH, 222-3°; Me(CH2)4NH, 215-16°;
BuCHMeNH, 186.5-8.5°; Et2CHNH, 209-11°; Me(CH2)5NH,
194.5-6.5°; cyclopropylmethylamino, 220-1.5°;
cyclopropylamino, 213-15°; cyclopentylamino, 219-20°;
PhCH2NH, 206-9°; p-MeC6H4CH2NH, 216-17°; o-FC6H4CH2NH,
206-8°; p-C1C6H4CH2NH, 225-6°; PhCH2CH2NH, - (HCl salt m.
199-202°); F3CCH2NH, 232-3°; F3CCH2CH2NH, 221-2.5°;
HOCH2CH2NH, - (HCl salt m. 272-3°); HOCH2(CHOH)4CH2NH,
223-4°; H2NCH2CH2NH, - (HCl salt m. 311°); Me2NCH2CH2NH,
192.5-4.5°; 4-pyridylmethylamino, 239-40°;
2-furylmethylamino, 217-18°; PhNH, 246.5-8.5°; p-ClC6H4NH,
276-8°; MeEtN, 229-3°; MeBuN, 214-15°; iso-PrMeN,
207-8°; Me(CH2:CHCH2)N, 207-8°; MeBuN, 208-9°; Et2N,
215°; EtPrN, 224-5°; iso-PrEtN, 207-8°;
Et(CH2:CHCH2)N, 208-9°; EtBuN, 200.5-1.5°; Pr2N,
221-2°; PrBuN, 215-17°; 1-pyrrolidinyl, 244.5-5.5°;
hexamethylenimino, 224-5°; 4-methylpiperazino, - (2HCl salt m;
229-300°); MeNHNH, 234°; Cl2N, - (HCl salt m.
259-61°); MeNH, 218-19° (decomposition); Me2NNMe, - [2HC1 salt m.
262° (decomposition)]; MeNH, 210° (decomposition) [sic]; Me2N,
245° (decomposition); MeBrN. - [HCl salt m. 288° (decomposition)];
EtNH, 207.5-9.5° (decomposition); cyclohexylamino, 221-2°
(decomposition); cycloheptylamino, 228-30° (decomposition); cyclopropylamino,
196.5-9° (decomposition); PhNH, 224-6° (decomposition); PhNH,
194.5-5.5° (decomposition)[sic]; Ph2N, 234.5-5.5°; PhClN,
214-16° (decomposition); PhBrN, 234-6° (decomposition); p-C1C6H4NH,
282-5° (decomposition); MePhN, 212-13° (decomposition); MePhN,
218-19° (decomposition)[sic]; Me2NNPh, 204-6° (decomposition);
1-pyrrolidinyl, 220-1°; 1-pyrryl, 211-13°;
3-chloro-1-pyrrolyl, 246-7° (decomposition); (3-isopropylidineamino-6-
anilino-2-pyrazinecarbonyl) guanidine, 214-16° (decomposition);
(3-acetoamido-6-methylthio-2-pyrazinecarbonyl)guanidine, 220-2°;
the following I (X = NH2, Y = Cl) (R, R1, m.p., and m.p. HCl salt given): H, HOCH2CH2, -, 228.5-9.5° (decomposition); H, Ph, -, -, [MeSO3H salt m.
272° (decomposition)]; H, PhCH2, 215-16° (decomposition); -; H,
p-FC6H4CH2, 216-19.5° (decomposition), -; H, PhCHMe, 153-60°
(decomposition), -; H, 2-C10H7CH2, 243.5-5.5° (decomposition), -; H,
3-pyridylmethyl, 280.5-3.5° (decomposition), -; H, p-MeC6H4CH2,
210-12° (decomposition), -; Me, PhCH2, 274.5° (decomposition), -; H,
o-C1C6H4CH2, 220-3° (decomposition), -; H, p-C1C6H4CH2, 204-6°
(decomposition), -; H, p-MeOC6H4CH2, 175.5-9.5° (decomposition), -; H,
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2,4-Me2C6H3CH2, 220-2° (decomposition), -; H, 2,4-C12C6H3CH2, -, 267.5-70.5° (decomposition); H, 3,4-C12C6H3CH2, 216-19° (decomposition), -; H, PhClH,CH2, 219-21° (decomposition), -; Me, Me, 240° (decomposition), -, [HCl.H2O salt m. 275° (decomposition)]; H, octahydrol-azocinyl, -, -; Et, Et, 265° (decomposition), -; Bu, Bu, 148-9°, -; (RR1 =) (CH2)4, -, -; (RR1 =) 3-oxapentamethylene, -, -; the following I (R = R1 = Me, Y = C1) (X and m.p. given): iso-PrNH, 238-40.5°; CH2:CHCH2NH, 213-15°; BuNH, 187.5°; cyclopropylmethylamino, 196-7°; Me2N, 219°; MeEtN, 217-18°; iso-PrMeN, 209-11°; Et2N, 212-14°; I (R = H, R1 = HOCH2CH2, X = iso-PrNH, Y = C1).HC1.0.5H2O [m. 185-6° (decomposition)], and 1-(3,5-diamino-6-chloro-2-pyrazinecarbonyl)2,3dimethylquanidine. 1634-20-4P RL: SPN (Synthetic preparation); PREP (Preparation)

(preparation of)

1634-20-4 CAPLUS RN

ΤТ

CN Pyrazinecarboxamide, N-amidino-3-amino-5,6-diphenyl- (7CI, 8CI) (CA INDEX NAME)

ANSWER 28 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1967:500105 CAPLUS DOCUMENT NUMBER: 67:100105

ORIGINAL REFERENCE NO.: 67:18835a,18838a

TITLE: Pyrazine diuretics. III. 5- and 6-alkyl, -cyclo-alkyl, and -aryl derivatives of

N-amidino-3-aminopyrazinecarboxamides AUTHOR(S): Bicking, John B.; Robb, Charles M.; Kwong, Sara F.;

Craque, Edward J., Jr.

CORPORATE SOURCE: Merck and Co. Inc., West Point, PA, USA

SOURCE: Journal of Medicinal Chemistry (1967), 10(4), 598-602

CODEN: JMCMAR: ISSN: 0022-2623

DOCUMENT TYPE: Journal LANGUAGE: English

For diagram(s), see printed CA Issue.

AB cf. CA 63: 11561e; 66: 37887h. In evaluations of N-amidino-3aminopyrazinecarboxamides as diuretics, a series of 5- and 6-alkyl, -cycloalkyl, and -aryl derivs. was synthesized and studied for effects on renal electrolyte excretion. Several compds. reverse the electrolyte excretion effects of deoxycorticosterone acetate in the adrenalectomized rat, the most highly active being N-amidino-3-amino-6methylpyrazinecarboxamide (I). 16 references.

1634-20-4P

RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

RN 1634-20-4 CAPLUS

CN Pyrazinecarboxamide, N-amidino-3-amino-5,6-diphenyl- (7CI, 8CI) (CA INDEX NAME)

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NH
C-NH-C-NH2
NHo
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L7 ANSWER 29 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1967:37882 CAPLUS DOCUMENT NUMBER: 66:37882

ORIGINAL REFERENCE NO.: 66:7227a,7230a

TITLE: Synthesis of furan derivatives. XXXIV. Preparation

of 2,3-bis(5-nitro-2-furyl)pyrazine derivatives AUTHOR(S): Saikachi, Haruo; Matsuo, Junro

CORPORATE SOURCE: Kyushu Univ., Fukuoka, Japan

SOURCE: Yakugaku Zasshi (1966), 86(10), 927-32 CODEN: YKKZAJ; ISSN: 0031-6903

DOCUMENT TYPE: Journal

LANGUAGE: Japanese

For diagram(s), see printed CA Issue.

cf. CA 61, 5648a. To a warm (40°) suspension of 19 g. furil in EtOH is added 8 g. ethylenediamine and the whole stirred 2 hrs. to give 20 g. 2,3-di(2-fury1)-5,6-dihydropyrazine (I), m. 128° (dilute EtOH). Similarly prepared is the 5-Me derivative of I, m. 94°, in 70% yield. (21 g.) is refluxed with 27 g. KCN in 250 ml. 80% EtOH for 30 min., the mixture filtered hot, and 3 vols. H2O is added to the filtrate to give 12 g. 5,6-di(2-furyl)-2-pyrazine-carboxamide (II), yellow plates, m. 182 (EtOH); similarly is prepared the 3-Me derivative, yellow plates, m. 175°. I (21 g.) in 250 ml. 80% EtOH is treated with 4 g. NaOH under introduction of air and 4 vols. H2O added to give 18 g. 2,3-di(2-furyl)pyrazine (III), yellow flakes, m. 81° (EtOH); similarly is prepared the 5-Me derivative, yellow flakes, m. 65°. III (4.2 g.) is dropped into a cold (-10°) mixture of 7.8 g. fuming HNO3 and 18 g. Ac20, the whole is made to react for 3 hrs., and poured into iced H2O to give 2 g. 2,3-bis(5-nitro-2-furyl)pyrazine, yellow prisms, m. 237° (dioxane); similarly prepared is the 5-Me derivative, yellow plates, m. 197° (AcOH). II (13 g.) is hydrolyzed with 10 g. NaOH in 300 ml. 50% EtOH to give 5,6-di(2-furyl)-2-pyrazine-carboxylic acid (IV), m. 151° (EtOH), almost quant.; the 3-Me derivative, yellow needles, m. 129°. IV (13.5 g.) is esterfied with 300 ml. EtOH and 10 g. concentrated H2SO4 to give 11 g. Et 5,6-di(2-furyl)-2-pyrazinecarboxylate (V), m. 98° (EtOH); the 3-Me derivative, m. 95°. V (2.8 g.) is gradually added to a cold (-5 to -10°) mixture of 3.9 g. fuming HNO3 and 9 g. Ac20, the whole stirred at the same temperature for 2 hrs., and poured into iced H2O to give 1.2 g. Et 5,6-bis(5-nitro-2-fury1)-2pyrazinecarboxylate (VI), m. 159° (AcOEt); the 3-Me derivative, yellow plates, m. 134°. VI (1.8 g.) is refluxed in a mixture of 100 ml. 50% AcOH and 2 ml. concentrated H2SO4 for 5 hrs. to give 1.5 g. 5,6-bis(5-nitro-2furv1)-2-pyrazinecarboxylic acid monohydrate, vellow prisms, m. 197° (EtOH); the 3-Me derivative, yellow needles, m. 206°. (decomposition). Also prepared are the VII tabulated. [TABLE OMITTED] 13480-81-4P 13484-30-5P 13484-31-6P

13484-35-0P 14399-30-5P 15541-91-0P RL: SPN (Synthetic preparation); PREP (Preparation) (preparation of)

13480-81-4 CAPLUS RN

Pyrazinecarboxamide, 5,6-di-2-furyl-3-methyl- (8CI) (CA INDEX NAME)

RN 13484-30-5 CAPLUS

CN Pyrazinecarboxylic acid, 3-methyl-5,6-bis(5-nitro-2-furyl)-, hydrazide (8CI) (CA INDEX NAME)

RN 13484-31-6 CAPLUS

CN Pyrazinecarboxanilide, 3-methyl-5,6-bis(5-nitro-2-furyl)- (8CI) (CA INDEX NAME)

RN 13484-35-0 CAPLUS

CN Pyrazinecarboxylic acid, 5,6-di-2-furyl-3-methyl-, hydrazide (8CI) (CA INDEX NAME)

14399-30-5 CAPLUS

Pyrazinecarboxamide, 3-methyl-5,6-bis(5-nitro-2-furyl)- (8CI) (CA INDEX CN NAME)

RN 15541-91-0 CAPLUS

CN Pyrazinecarboxanilide, 4'-hydroxy-3-methyl-5,6-bis(5-nitro-2-furyl)- (8CI) (CA INDEX NAME)

ANSWER 30 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

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INVENTOR(S):

Cragoe, Edward J., Jr. Merck & Co., Inc.

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A suspension of 765 g. Me 3-aminopyrazinecarboxylate in 5 l. C6H6 was treated with 1.99 1. SO2C12, refluxed for 5 hrs., and left overnight at room temperature to give 888 g. crude Me

3-amino-5,6-dichloropyrazinecarboxylate

(I), m. 233-4°. Into a solution of 100 g. I in 1 l. dry Me2SO dry NH3 was passed under stirring at 65-70° for 45 min., then at 10° for 1.25 hrs. to give 82.5 g. Me 3,5-diamino-6-chloropyrazinecarboxylate (II), m. 212-13°. A mixture of 14.2 g. II, 9 g. Pd-C, 4 g. MgO, and 250 ml. MeOH was shaken under H for 18 hrs. at room temperature to give Me 3,5-diaminopyrazinecarboxylate (III), m. 252-4° (decomposition) (iso-PrOH). Bromination of a suspension of 2 g. III in 25 ml. AcOH at 50° with 2.1 g. Br in 10 ml. AcOH gave 1.2 g. Me 3,5-diamino-6-bromopyrazinecarboxylate (IV), m. 217-19°. Hg(OAc)2 (3.2 g.) and a solution of 2.5 g. iodine in 20 ml. warm dioxane was added rapidly to a suspension of 1.7 q. III in 30 ml. H2O at 70°, the mixture heated for 5 min., cooled to room temperature, and treated with 50 ml.

15%

KI solution precipitated 1.2 g. Me 3,5-di-amino-6-iodopyrazinecarboxylate, m. 200-2°. I (11.1 g.), 500 ml. iso-PrOH, 14.4 g. PhNH2, and 12.8 g. PhNH2.HCl was refluxed 24 hrs. under stirring to give 10 g. Me 3-amino-5-anilino-6-chloropyrazinecarboxylate, m. 171.5-73° (iso-PrOH). Similarly were prepared Me 3-amino-5-(p-chloroanilino)-6chloropyrazinecarboxylate, m. 207-8° (MeCN), and Me 3-amino-5-dimethylamino-6-chloropyrazinecarboxylate (V), m. 145.5-6.5° (MeOH). A solution of 10 g. MeSH in 17 ml. 20% NaOH and 100 ml. MeOH was added to a boiling mixture of 17.7 g. I and 1 l. MeOH and refluxed 15 min. to precipitate 12 q. Me 3-amino-5-methylthio-6chloropyrazinecarboxylate (VI), m. 212-16° (MeOH). VI (23.4 g.), 35 ml. 30% H2O2, and 300 ml. AcOH was stirred 18 hrs. at room temperature to give 18.5 g. the 5-methylsulfinyl analog (VII), m. 237.5-40.5° (decomposition) (MeOH-AcOEt-HCONH2). Hydrolysis of 7.5 g. VII in 75 ml. AcOH and 12 ml. H2O on a steam bath for 3 hrs. produced 3.7 g. Me 3-amino-5-hydroxy-6-chloropyrazinecarboxylate (VIII), m. .apprx.245° (decomposition) (HCONH2-EtOH). Hydrogenation of VIII with Pd-C and MgO at room temperature resulted in Me 3-amino-5hydroxypyrazinecarboxylate, decompose 220-60°. Also were prepared Me 3-amino-5-dimethyl-aminopyrazinecarboxylate, m. 242.5-3.5°, Me 3,5-diaminopyrazinecarboxylate, m. 252-4° (decomposition), and Me 3-amino-5-methoxypyrazinecarboxylate, m. 205.5-7.5°. A mixture of 8.9 q. I and 20 ml. PhCH2NH2 was heated on a steam bath for 30 sec. to give 7.5 g. Me 3-amino-5-benzylamino-6-chloropyrazinecarboxylate (IX), m. 157-8° (MeOH). Hydrogenation of IX yielded Me 3-amino-5benzylaminopyrazinecarboxylate, m. 189.5-91.5°. Treatment of 1.1 g. I with MeONa in 200 ml. boiling absolute MeOH produced 1 g. Me 3-amino-5-methoxy-6-chloropyrazinecarboxylate, m. 255-7° (MeCN). Na2S (9.6 g.) and 10 g. S was refluxed in 80 ml. absolute EtOH. Addition of g. I at 25° and stirring for 1 hr. gave 7.8 g. Me

8.9 (decomposition). To a refluxing solution of 4.44 g. I in 300 mil EtOH was

added quanidine (from 1.98 g. quanidine-HCl) in 50 ml. absolute EtOH in 15 min. and

3-amino-5-mercapto-6-chloropyrazinecarboxylate, m. 207-8°

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the mixture refluxed 0.5 hr. to give 3.1 g. Me 3-amino-5-ethoxy-6-
     chloropyrazinecarboxylate, m. 123-5° (iso-PrOH).
     3-Amino-6-methylpyrazinoylamide (31 g.) was heated 10 min. with 320 ml.
     10% NaOH. The resulting Na salt of the acid (97 g.) was methylated with
     77 g. Me2SO4 in 700 ml. MeOH 19 hrs. at room temperature to give 18 g. Me
     3-amino-6-methylpyrazinecarboxylate (X), m. 138.5-40.5° (C6H6).
     Chlorination of 9.2 q. X with 65 ml. SO2C12 under cooling produced 4.4 q.
     Me 3-amino-5-chloro-6-methylpyrazinecarboxylate, m. 108.5-10.5°
     (C6H6-cvclohexane). A mixture of 30 g. 3-amino-5-methylpyrazinecarboxylic
     acid and a solution of 30% HCl in 650 ml. MeOH was stirred 42 hrs. at room
     temperature to give 15.4 g. Me 3-amino-5-methylpyrazinecarboxylate (XI), m.
     165-7° (H2O). A solution of 4.18 g. Br in 3 ml. AcOH was added to a
     solution of 4.2 g. XI in 15 ml. AcOH in 20 min. to produce 3.6 g. Me
     3-amino-5-methyl-6-bromopyrazinecarboxylate, m. 179-81°.
     Aminomalonamidamidine-2HCl (52.5 g.) was added to an ice-cooled solution of
     28.8 g. ethylglyoxal in 450 ml. H2O. The mixture was made alkaline with
     .apprx.65 ml. concentrated NH4OH and left 20 hrs. at room temperature to
precipitate 17.5 g.
     3-amino-6-ethylpyrazinecarboxamide, m. 165.5-8.5° (iso-PrOH), which
     was saponified 30 min. on a steam bath with 10% NaOH to give
     3-amino-6-ethylpyrazine-carboxylic acid (XII), m. 149-52°.
     Stirring 14 g. XII in a solution of 33% HCl in 160 ml. MeOH 24 hrs. at room
     temperature gave 4.3 g. XII Me ester, m. 85-7° (iso-PrOH). Also prepared
     were 3-amino-6-p-chlorophenylpyrazinecarboxylic acid, m. 207-13°,
     and its Me ester, m. 181.5-3.5°. To a suspension of 17.9 g.
     5,6-diaminouracil in 250 ml. H2O at 60° 14.9 q.
     cyclohexylgiyoxal-0.5 H2O was added and the mixture heated 1 hr. on a steam
     bath to give 7.5 g. 7-cyclohexyllumazine (XIII), m. 229-31° (aqueous
     AcOH). A solution of 18.5 g. XIII and 9 g. NaOH in 90 ml. H2O was heated in
     an autoclave 17 hrs. at 105° to give 8 g. 3-amino-5-
     cyclohexylpyrazinecarboxylic acid, m. 182.5-3.5° (aqueous iso-PrOH); Me
     ester m. 173-4.5°. Similarly were prepared Me 3-amino-6-
     cyclohexylpyrazinecarboxylate, m. 126.5-28°, Me
     3-amino-6-cyclopropylpyrazinecarboxylate, m. 112.5-14.5° (amide m.
     185.5-7.5°, free acid m. 169-72°), Me 3-amino-5-
     phenylpyrazinecarboxylate (XIV), m. 231-2°, and Me
     3-amino-6-phenylpyrazinecarboxylate (XV), m. 140-1°. Chlorination
     of 25.6 g. XV with 90 ml. SO2Cl2 1.5 hrs. at room temperature gave Me 3-amino
     5-chloro-6-phenylpyrazinecarboxylate, m. 187.5-91.5° (AcOH).
     Bromination of 10.5 q. XIV in 700 ml. AcOH with 11.2 q. Br 21 hrs. at
     85° gave 10.5 g. Me 3-amino-5-phenyl-6-bromopyrazinecarboxylate, m.
     217-21° (AcOH). To a suspension of 103.59 g. 4,5-diamino-2,6-
     dihydroxypyrimidine in 1500 ml. H2O and 500 ml. concentrated NH4OH at 60°
     103.71 g. 1-phenyl-1, 2-propanedione was added and the mixture heated at
     90° under vigorous stirring to give 82.4 g. 6(or 7)-methyl-7(or
     6)-phenyllumazine, m. 281.5-2.5° (AcOH), and 32 g. 6(or
     7)-phenyl-7(or 6)-methyllumazine (XVI), m. 254.5-5.5°. Saponification of
     XVI with 8% NaOH in an autoclave 3.5 hrs. at 170° gave 3-amino-5(or
     6)-phenyl-6(or 5)-methylpyrazinecarboxylic acid, m. 193.5-4.5°; Me
     ester m. 163-4° (MeOH). Similarly were prepared 3-amino-5(or
     6)-methyl-6(or 5)-phenylpyrazine carboxylic acid, m. 155-6°; Me
     ester m. 162.5-3.5° (MeOH). Me 3-amino-6-phenylpyrazinecarboxylate
     was chlorinated with SO2C12 to give Me 3-amino-5-chloro-6-
     phenylpyrazinecarboxylate, m. 187.5-90.5° (AcOH), and subsequently
     treated with Me2NH in MeOH to give Me 3-amino-5-dimethylamino-6-
     phenylpyrazinecarboxylate, m. 167.5-9.5° (MeOH). To 750 ml. AcOH
     and 3180 ml. H2O at 38°, 90 g. Me 3-aminopyrazinecarboxylate was
     added and Cl passed through in 25 min. to give Me 3-amino-6-
     chloropyrazinecarboxylate (XVII) m. 142° (decomposition) (H2O). A solution
     of 18.8 g. XVII, 15 g. PhNH2, and 2.5 ml. concentrated HCl in 150 ml. Me2CO was
     refluxed 16 hrs. to give 7.4 g. Me 3-isopropylideneamino-6-
     anilinopyrazinecarboxylate, m. 195.5-7.5° (iso-PrOH). A mixture of
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9.3 q. 3-amino-5,6,7,8-tetrahydroquinoxaline-2-carboxylic acid and 230 ml.
     absolute MeOH of 10° was treated with 30 ml. concentrated H2SO4 in 1 hr. and
     left 24 hrs. at room temperature to give 1.6 g. the Me ester, m. 154-5°
     (1:5 MeOH-H2O). A solution of 60 g. 4-chloro-o-phenylenediamine in
     60 ml. H2O and 50 ml. 12N HCl was treated with a solution of 61.44 g.
     alloxan-H2O in 100 ml. H2O and stirred 1 hr. at 90° to give a precipitate
     of 78.4 g. 8-chloroalloxazine, m. 365-6° and 40.36 g.
     7-chloro-alloxazine, (XVIII) m. 380° (Me2SO). A mixture of 44.2 q.
     XVIII and 190 ml. concentrated NH4OH was heated in an autoclave 10 hrs. at
     165° to give 27.2% 3 amino-7-chloroguinoxalin-2-carboxylic acid, m.
     191-2° (decomposition); Me ester m. 224.5-5.5° (MeCN). Also
     prepared are the following XIX (R, R1, % yield, and m.p. given): Me, H, 88,
     221-2°; Et, H, 89, 149-50°; Pr, H, 75, 138-40°;
     iso-Pr, H, 70, 125.5-6.5°; CH2:CHCH2, H, 69, 105-6.5°; Bu,
     H, 91, 140-2°; sec-Bu, H, 75, 106-8°; iso-Bu, H, 51,
     113.5-15.5°; tert-Bu, H, 38, 98-108°; Am, H, 72,
     100.5-2.5°; MePrCH, H, --, --; Et2CH, H, --, --; C6H13, H, 70,
     72.5-5.5°; cyclopropylnethyl, H, 78, 132-3° cyclopropyl, H,
     98, 167-9°; cyclopentyl, H, 93, 119.5-21.5°; PhCH2, H, 64,
     157-8°; p-MeC6H4CH2, H, 66, 112.5-14.5°; o-FC6H4CH2,
     H, 84, 171-4°; p-C1C6H4CH2, H, 93, 136-7°; PhCH2CH2, H, 59,
     115-19°; CF3CH2, H, 97, 153-4° CF3CH2CH2, H, 76, 124.5-5.5°; HOCH2CH2, H, 100, 155-7°; HOCH2(CH0H)4CH2, H,
     60, 172-5°; NH2CH2CH2, H, 96, 265°; Me2NCH2CH2, H, 40,
     257°; 4-pyridylmethyl, H, 69, 95-7°; 2-furylmethyl, H, 81,
     148-9°; Me, Et, 73, 102-4°; Me, Pr, 58, 83.5-5.5°; Me, iso-Pr, 78, 75.5-7.5°; Me, CH2:CHCH2, 70, 90.5-92°; Me,
     Bu, 74, 59.5-61.5°; Et, Et, 54, 99-101°; Et, Pr, --, --; Et,
     iso-Pr, --, --; Et, CH2:CHCH2, --, --; Et, Bu, 91, 77.5-9.5°; Pr,
     Bu, --, --; Pr, Pr, 66, 68.5-71.5°; (NRR1 = ) pyrrolidino, 95,
     168-71°; (NRR1 =) 1 (hexahydroazepiny1), 75, 109-11°; (NRR1
     N'-Methylpiperazino, 88, 186-8°; Me, NH2, 67, 136.5-38°
     Guanidine-HCl (XX) (26.3 g.) was added to a solution of MeONa (5.75 g. Na in
     150 ml. absolute MeOH), the precipitated NaCl filtered off, and the filtrate
concentrated
     to 30 ml. After addition of 11.5 g. V the mixture was boiled 1 min., then
     maintained 1 hr. at room temperature to give 93% (3-amino-5-dimethylamino-6-
     chloropyrazinecarbonyl) quanidine (XXa), m. 216-17°; HCl salt m.
     298° (decomposition). Similarly were prepared (3,5-diamino-6-bromopyrazin-
     carbonyl)quanidine, m. 232.5-5.5° (decomposition), (3,5-diamino-6-
     iodopyrazinecarbonyl)quanidine-HCl, m. 273-4° (decomposition) and
     (3-isopropylideneamino-6-anilinopyrazinecarbonyl) quanidine, m.
     214-16° (decomposition). To a solution of 920 mg. Na in 50 ml. absolute
     iso-PrOH 3.85 g. XX was added and the NaCl filtered off. Adding 4.4 g. I
     and refluxing the mixture 15 min. gave (3-amino-5,6-
     dichloropyrazinecarbonyl)quanidine HCl salt (XXb) m. 259-61°. The
     solution of XXb in 5 ml. HCONMe2 was treated with 1 ml. 25% aqueous Me2NH 1 hr.
     on a steam bath to give XXa. Reaction of 11.1 g. I with 55 ml.
     Me2NCH2CH2OH 20 min. on a steam bath gave 9.5 g. Me 3-amino-5-(2-
     dimethylamino-ethoxy)-6-chloropyrazinecarboxylate (XXI), m.
     134.5-6.5° (C6H6-cyclohexane). To 20 g. XX in iso-PrONa (4 g. Na in 100 ml. iso-PrOH) 9.4 g. XXI was added and the mixture heated 30 min. on
     a steam bath to give 2.5 g. (3-amino-5-guanidino-6-
     chloropyrazinecarbonyl)guanidine-2HCl, m. >340^\circ. A mixture of 2 1. concentrated NH4OH and 300 g. XVIII was stirred 16 hrs. at room temperature to
give
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260 g. 3-amino-6-chloropyrazinecarboxamide (XXII), m. 227-30°. HC(OEt)3 (200 ml.) and 33 g. XXII refluxed in 200 ml. Ac20 1.5 hrs. gave 20 g. 4-hydroxy-6-chloropteridine (XXIII), m. 268-70° (decomposition) (iso-PrOH). A solution of 5.5 g. XXIII and 4.4 g. PhCH2SH in 100 ml. 4% NaOH was heated 30 min. on a steam bath to give 5.5 g. 4-hydroxy-6-benzylthiopteridine, m. 233-5° (aqueous iso-PrOH), which was converted

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into 3-amino-6-benzylthiopyrazinecarboxylic acid (XXIV), m. 138-9°,
by 8 hrs. hydrolysis with 5% NaOH. XXIV (8.5 q.) in 50 ml. Ac20 was
heated 5 hrs. on a steam bath to give 6.6 g. 2-methy1-6-benzy1thio-4H-
pyrazino[2,3-d][1,3]oxazin-4-one (XXV), m. 116.5-18.5° (C6H6). To
1 g. Na in 30 ml. iso-PrOH 5 g. XX and 3.4 g. XXV were added to give,
after 1 hr. at room temperature, 1.1 g. (3-amino-6-benzylthiopyrazinecarbonyl-
quanidine, m. 171-3° (decomposition). Similarly were prepared
4-hydroxy-6-methylthiopteridine, m. 289.5-91.5° (aqueous iso-PrOH),
3-amino-6-methylthiopyrazinecarboxylic acid (XXVI), m. 182-4°
(decomposition) (AcOEt), 2-methyl-6-methylthio-4H-pyrazino[2,3-d][1,3]oxazin-4-
one, m. 189-91° (C6H6), and 3-acetamido-6-
methylthiopyrazinecarbonyl)quanidine (XXVII), m. 220-2°. Addition of
HCl to XXVII in H2O gave 86% (3-amino-6-methyl-
thiopyrazinecarbonyl)guanidine, m. 203-5°. A solution of 0.92 g. XXVI
in 15 ml. 2.5% NaOH was treated with 1.05 g. KMnO4 in 35 ml. H2O to give
0.5 g. 3-amino-6-methylsulfonylpyrazine-carboxylic acid, m. 239-42°
(decomposition) (iso-PrOH), which gave, after 5 hrs. heating in Ac20,
2-methyl-6-methylsulfonyl-4H-pyrazino[2,3-d][1,3]oxazin4-one, m.
214-16° (Me2CO), transformed into 27% 3-amino-6-
methylsulfonylpyrazinecarbonyl)guanidine, m. 224-6° (decomposition)
(iso-PrOH). Similarly are prepared the following XXVIIa (R, R1, % yield,
and m.p. given): H, H, 93, 240.5-1.5°; 293.5° (HCl salt);
Me, H, 89, 238-9°; Et, H, 63, 217-18°; Pr, H, 93,221-2°; iso-Pr, H, 75, 215°; CH2:CHCH2, H, 84,
213-14°; Bu, H, 65, 219.5°; Me-ETCH, H, 74, 208-9°;
iso-Bu, H, 76, 221°; tert-Bu, H, 84, 222-3°; Am, H, 70, 215-16°; MePrCH, H, 89, 186.5-8.5°; Et2CH, H, 82, 209-11°; C6H13, H, 100, 194.5-6.5°; cyclopropylmethyl, H,
95, 220-1°; cyclopropyl, H, 85, 213-15°; cyclopentyl, H
65, 219-20°; PhCH2, H, 44, 206-9°; p-MeC6H4CH2, H, 57,
216-17°; o-FC6H4CH2, H, 100, 206-8°; p-C1C6H4CH2,
H, 96, 225-6°; PhCH2CH2, H, 57, 199-202°; CF3CH2, H, 77,
232-3°; CF3CH2CH2, H, 65, 221-2.5°; HO-CH2CH2, H, 63,
272-3°; HOCH2 (CHOH) 4CH2, H, 68, 223-4°; NH2CH2CH2, H, 68,
311°; Me2NCH2CH2, H, 98, 192.4-4.5°; 4-pyridylmethyl, H, 64,
239-40°; o-furylmethyl, H, 92, 217-18°; Ph, H, 95,
246.5-8.5°; p-C1C6H4, H, 95, 276-8°; Me, Et, 92,
229-30°; Me, Pr, 97, 214-15°; Me, iso-Pr, 70, 207-8°;
Me, CH2:CHCH2, 95, 207-8°; Me, Bu, 95, 208-9°; Et, Et, 75,
215°; Et, Pr, 92, 224-5°; Et, iso-Pr, 75, 207-8°; Et,
CH2:CHCH2, 92, 208-9°; Et, Bu, 98, 200.5-1.5°; Pr, Pr, 100,
221-2°; Pr, Bu, 84, 215-17°; (NRR1 =) pyrrolidino, 90,
244.5-5.5°; (NRR1 =) 1-hexahydroazepiny1, 49, 224-5°; (NRR1
N-methylpiperazino, 74, 299-300°; Me, NH2, 92, 234°.
Also prepared are the following XXVIIb (X, Y, % yield, and m.p. base and
m.p. HCl salt given): H, HO, 10, >310° (decomposition); H, NH2, 8,
286-8° (decomposition), --; H, NMe2, 45, 224-5° (decomposition), --;
H. MeO, 52, --, 229-30° (decomposition); H. PhCH2NH, 56, --,
231-7° (decomposition); Cl, MeO, 90, --, 257°; Cl, MeS, 100,
234.5-6.5°, --; Cl, HO, 24, --, >300° (decomposition); Cl, SH, 100, 236.5° --; Cl, EtO, 81, 215-16° --; Cl, Cl, 72, --,
259-61°; Me, H, 87, 218-19 (decomposition), --; Me, Me2N, 42, --, 262° (decomposition) (di-HCl); H, Me, 13,210° (decomposition), --; Me,
Me, 38, 245° (decomposition), --; Br, Me, 35, 288° (decomposition),
--; Et, H, 53, 207.5-9.5° (decomposition), --; H, cyclohexyl, 71,
221-2° (decomposition), --; cycloheptyl, H, 61, 228-30°
(decomposition), --; cyclopropyl, H, 61, 196.5-99° (decomposition), --; H,
Ph, 51, 224-6° (decomposition); Ph, H, 34, 194.5-5.5° (decomposition),
--; Ph, Ph, 87, 234.5-5.5°, -; Ph, Cl, 69, 214-16°
(decomposition), --; Br, Ph, 66, 234-6° (decomposition), --; p-ClC6H4, H, 70,
282-5° (decomposition), --; Me (or Ph), Ph (or Me), 77, 212-13°
(decomposition), --; Ph (or Me), Me (or Ph) 90, 218-19° (decomposition), --;
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Ph, Me2N, 40, 205-6° (decomposition), --; (XY =) (CH2) 4, 29,
     220-1°, --; (XY =) CH:CHCH:CH, 56, 211-13°, --; (XY =)
     HC:CC1CH:CH, 70, 246-7° (decomposition), --. A solution of 13.9 g.
     2-methyl-2-pseudothiuronium sulfate (XXVIII) and 9.2 g. H2NCH2CH2OH in 40
     ml. H2O was heated 20 min. to give 12.5 g. (2-hydroxyethyl)guanidine
     sulfate, m. 127.5-35.5°, which was added to a solution of 2g. Na in 25
     ml. MeOH, MeOH distilled, and the residue treated with 4.1 q. II 5 min. on
     steam bath to give 1.2 g. 1-(3,5-diamino-6-chloropyrazinoy1)-3-(2-
     hydroxyethyl)quanidine-HCl, m. 228.5-9.5° (aqueous iso-PrOH).
     1-(3-Amino-5-isopropylamino-6-chloropyrazinov1)-3-(2-
     hydroxyethyl)guanidine-HCl.0.5H2O, m. 185-6° (decomposition), was prepared
     from Me 3-amino-5-isopropylamino-6-chloropyrazinecarboxylate. A mixture of
     6.1 g. II, 6.8 g. phenylquanidine, and 3 ml. iso-PrOH was heated 6 hrs. to
     give 1-(3.5-diamino-6-chloropyrazinoy1)-3-phenylguanidine, isolated as the
     MeSO3H salt, m. 272° (decomposition) (H2O). Ph-CH2NH2 (80.3 g.) and
     69.5 g. XXVIII in 200 ml. H2O kept 18 hrs. at room temperature gave
     benzylguanidine sulfate, which was converted into the HCl salt (XXIX)
     (51.5 g.), m. 175-8° (aqueous EtOH), by treating its aqueous solution with
aqueous
     BaCl2. To a solution of 1 g. Na in 30 ml. iso-PrOH 9.3 g. XXIX was added and
     half the volume distilled Addition of 2 g. II and heating the mixture 15 min.
     vielded 1 g. 1-(3,5-diamino-6-chloropyrazinov1)-3-benzylguanidine, m.
     215-16° (decomposition) (aqueous iso-PrOH). With the appropriate starting materials the following 3-substituted 1-(3,5-diamino-6-
     chloropyrazinoyl) quanidines were prepared [3-substituent and m.p.
(decomposition)
    given]: p-fluorobenzyl 216-19.5°; a-methylbenzyl
     153-60°; 3-pyridylmethyl, 280.5-3.5°; 2-naphthylmethyl
    243.5-5.5°. Also prepared were the following RR1-NC(:NH)NH2.HCl (R, R1, % yield, and m.p. given): p-Me-C6H4CH2 H, 28, 153-5°;
     o-C1C6H4CH2, Me, 32, 122.5-5.5°; PhCH2, H, 71,
     131-6°; p-C1C6H4CH2, H, 55, 162.5-4.5°; p-MeOC6H4CH2, H, 69,
     132-7°; 2,4-Me2C6H3CH2, H, 52, 105-15°; 2,4-C12C6H3CH2, H,
     67, 145-8°; 3,4-C12C6H4CH2, H, 77, 155-7°; PhCH2CH2, H, 71,
     135-8°.
  Also prepared were the following XXIXa [R, R1, % yield, and m.p.
     (decomposition)given]: p-MeC6H4CH2, H, 27, 210-12°; PhCH2, Me, 35,
     274.5° (HCl salt); o-ClC6H4CH2, H, 39, 220-3°;
     p-C1C6H4CH2, H, 46, 204-6° p-MeOC6H4CH2, H, 27, 175.5-9.5°;
     2,4-Me2C6H3CH2 H, 59, 220-2°; 2,4-C12C6H3CH2, H, 30,
     267.5-70.5° (HCl salt); 3,4-C12C6H3CH2, H, 47, 216-19°;
     PhCH2CH2, H, 46, 219-21.5°. To a solution of 2.3 g. Na in 200 ml.
    absolute MeOH 15 g. dimethyl-quanidine sulfate was added, the mixture refluxed
     hr. and cooled, Na2SO4 filtered off, the solution concd, to 30 ml., 10.15 g.
     II added, and the mixture heated 30 min. and kept 1 hr. at room temperature to
     give 3.6 g. 1-(3,5-diamino-6-chloropyrazinov1)-3,3-dimethyl-quanidine
     (XXX), decomposing at 240° HCl salt m. 275° (decomposition). To a
     solution of 36.57 g. Et2NH in 100 ml. H2O and 41 ml. concentrated HCl adjusted,
     with 3.66 g. Et2NH to pH 9.2 a solution of 50% aqueous cyanamide (65.16 g.) was
     added dropwise at 100° in 4 hrs. After refluxing 1 hr. and
     standing over night at room temperature the mixture was treated with 50 ml. of
40%
     NaOH and CO2 passed through under cooling to give 1,1-diethylguanidine,
     isolated as the HCl salt (XXXI) (35 g.), m. 147-9°. Similarly,
     1,1-dibutylquanidine-HCl (XXXII), m. 104.5-106° (H2O), was obtained
     in 86% yield. The following compds. were also prepared: 88.6% 1 -
     (3,5-diamino-6-chloropyrazinoy1)-3,3-diethylguanidine, m. 265°
     (decomposition), from II and XXXI and 72% 1-(3,5-diamino-6-chloropyrazinoy1)-
     3,3-dibutylguanidine, m. 148-9° (iso-PrOH), from II and XXXII.
    Also prepared were the following XXXIII (R, R1, % yield, and m.p. given):
     iso-Pr, H, 35, 238.5-40°; CH2:CHCH2, H, 39, 215°; Bu, H, 17,
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 $187.5^\circ;$ cyclopropylmethyl, H, 3, $196-7^\circ;$ Me, Me, 69, $219^\circ;$ Me, Et, 49, $218^\circ;$ Me, iso-Pr, 61, $209-11^\circ;$ Et, Et, $40,214^\circ.$ The compds. are effective in the treatment of abnormal electrolyte excretion.

IT 1634-20-4P, Pyrazinecarboxamide, N-amidino-3-amino-5,6-diphenyl-RL: PREP (Preparation)

(preparation of)

RN 1634-20-4 CAPLUS

CN Pyrazinecarboxamide, N-amidino-3-amino-5,6-diphenyl- (7CI, 8CI) (CA INDEX NAME)

DOCUMENT TYPE:

L7 ANSWER 31 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1961:38069 CAPLUS
DOCUMENT NUMBER: 55:38069

ORIGINAL REFERENCE NO.: 55:7423b-i,7424a-h

TITLE: Pteridines. XXIII. A facile pyrimidine ring cleavage
AUTHOR(S): Taylor, Edward C., Jr.; Knopf, Robert J.; Cogliano, J.

A.; Barton, J. W.; Pfleiderer, Wolfgang

CORPORATE SOURCE: Princeton Univ., Princeton, NJ SOURCE: Journal of the American Chemical Society (1960), 82,

6058-64

CODEN: JACSAT; ISSN: 0002-7863 Journal

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OTHER SOURCE(S): CASREACT 55:38069

cf. CA 55, 551q. 4-Mercaptopteridines and -pyrimidines were readily cleaved by C1CH2CO2H (I) and alkali carbonate or MeI and alkali. The results of a study of this cleavage indicated that heterocyclic systems containing a fused 4-substituted pyrimidine ring underwent a base-catalyzed cleavage to an o-aminonitrile, provided that the anion formed by attack of base at C-2 of the fused pyrimidine ring was capable of stabilization by appropriate structural features in the remainder of the mol., and that the substituent group attached to C-4 was capable of departure with its bonding pair of electrons in an irreversible cleavage step. These results underscored a fundamental chemical difference between purines and pteridines. 4-Mercapto-6,7-diphenylpteridine (0.2 g.) and 0.1 g. I in 15 cc. N NaHCO3 refluxed 0.5 hr. and filtered hot gave 0.12 g. 2-amino-3-cyano-5,6diphenylpyrazine (II), m. 160-3°; the aqueous phase from a similar run with a slight deficiency of Na2CO3 treated with AgNO3 gave the insol. Ag salt of HSCH2CO2H. II (0.54 q.), 0.16 q. NaOH, and 2 cc. 30% H2O2 in 25 cc. 40% aqueous EtOH refluxed 3 hrs. gave 0.40 g. 2-amino-5,6-diphenylpyrazine-3-carboxamide (III), yellow needles, m. 202-5°. II (1.4 g.) in 100 cc. 95% EtOH containing a few drops N(CH2CH2OH)3 treated 3 hrs. at 50-5° with H2S, the whole cooled, and filtered yielded 1.3 g. 3-CSNH2 analog of III, yellow needles, m. 158-60°. 4-Mercaptopteridine (IV) (0.5 g.), 0.45 g. I, 0.81 g. Na2CO3, and 30 cc. H2O refluxed 6 min., the mixture cooled to 0°, and filtered after 12 hrs. at 0° yielded 0.12 g. 2-amino-3-cyanopyrazine (V), needles, m. 192°; 0.04 g. 2nd crop. 4-MeS analog (VI) (0.54 g.) of V and 20 cc. N NaHCO3 refluxed 6 min., the mixture filtered, and the filtrate evaporated,

the residue sublimed at $150^\circ/0.5$ mm., and the sublimate $(0.2~\mathrm{g.})$ extracted with Et20 left $0.07~\mathrm{g.}^2$ -aminopyrazine-3-carboxamide (VII), needles, m. 235°, the residue from the Et20 extract recrystd. from H20 gave $0.09~\mathrm{g.}$ V, needles, m. 188- 90° ; the sublimation residue recrystd. from H20 gave a small amount of 4-hydroxypteridine (VIII). VI $(0.18~\mathrm{g.})$ and $10~\mathrm{cc.}$ N NaHCO3 refluxed $2~\mathrm{min.}$, the mixture filtered hot, and the filtrate cooled gave $0.1~\mathrm{g.}$ unchanged VI, m. 194° ; the filtrate contained V, VII, and VIII. VI $(0.16~\mathrm{g.})$ and $10~\mathrm{cc.}$ N NaHCO3 refluxed $45~\mathrm{min.}$ gave a mixture of VI, VIII, and 2-amino-3-carboxylic acid; the mixture evaporated, and the residue sublimed at $150^\circ/0.5~\mathrm{mm.}$ yielded $0.07~\mathrm{g.}$ VII, m. 230° . VI $(0.16~\mathrm{g.})$ and $10~\mathrm{cc.}$ N AcOH refluxed 1 hr. (MeSH evolved), the solution filtered hot with C_c and cooled to 0° yielded $0.1~\mathrm{g.}$ VIII. $\mathrm{HC}(0\mathrm{Et})3$ (60 cc.), $60~\mathrm{cc.}$ Ac2O, and $8.0~\mathrm{g.}$

about

1/3 of the original volume, diluted with 150 cc. dry Et2O, and cooled to 0° gave 6.30 g. 4-hydroxypyrimidol(4,5-d]pyrimidine (IX), needles, m. 253-5° (decomposition) (B2O). Powdered IX (3.70 g.) and 5.55 g. P2S5 in 20 cc. dry CSH5N refluxed 45 min., the mixture kept 15 min., poured with stirring into 50 cc. H2O and 50 g. crushed ice, stirred 0.5 hr., kept 12 hrs. at 0°, and filtered gave 3.80 g. 4-SH analog (X) of IX, bright yellow, did not melt but darkened rapidly above 300° (sublimed at 230°/0.1 mm.). X (0.66 g.) in 16 cc. 1% aqueous NaOH treated at 0-5° with 0.20 cc. MeI, the mixture stirred 1.5 hrs., filtered, and refrigerated overnight gave 0.40 g. 4-MeS analog (XI) of IX, m. 159-60° (sublimed at 130°/0.05 mm.). X (0.70 g.), 0.75 g. NaOH, and 12 cc. H2O stirred at room temperature to solution and then 2 hrs.

with

1.0 g. MeI, the whole cooled, and filtered gave 0.25 g. 4-amino-5-cyanopyrimidine, needles, m. 250-2° (H2O); also obtained in 82% yield by stirring XI in dilute aqueous NaOH at room temperature 4-Hydroxypyrid

o[3,4-d]pyrimidine (10 g.) and 59 g. P2S5 in 250 cc. dry C5H5N refluxed 2 hrs. and the solution evaporated in vacuo, the residue treated with 500 cc.

H20,

the mixture refluxed 20 min. after 12 hrs., and filtered, and the filter residue dissolved in 15 cc. H2O and 20 cc. concentrated NH4OH, the solution filtered, and added dropwise to 300 cc. refluxing H2O and 50 cc. AcOH gave 9.0 g. 4-mercaptopyrido[3,4-d]pyrimidine (XII) derivative of X, m. 325 (decomposition). XII (2.0 g.) in 20 cc. N NaOH and 10 cc. H2O shaken 5 min. with 1.5 cc. Me2SO4 and filtered gave 1.5 g. 4-MeS analog of XII. 4-Amionicotinic acid (XIII) (36 g.), 500 cc. absolute EtOH, and 36 cc.

concentrated

H2SO4 refluxed 70 hrs. on the steam bath and the whole worked up gave 31 g. Et ester (XIV) of XIII, m. 100-5°. XIV (25 g.) and 50 cc. HCONH2 heated 1 hr. at 160°, the mixture refluxed 3 hrs., cooled, and filtered vielded 10 q. 4-hydroxypyrido[4,3-d]pyrimidine (XV), m. 293° (H2O); 3.5 g. 2nd crop. XV was converted in the usual manner to the 4-SH analog (XVI) of XV, yellow, m. 323-5° (decomposition) (EtOH). XVI (1 g.), 0.9 g. I, 1.8 g. Na2CO3, and 30 cc. H2O refluxed 20 min., the mixture filtered, and cooled gave 0.15 g. 2-aminonicotinonitrile (XVII), m. 131°; the filtrate evaporated, and the residue sublimed at 120°/0.5 mm, gave 0.05 g. XVII; further sublimation at 200° yielded 0.1 g. 2-aminonicotinamide, m. 199°. XII (1 g.), 0.9 g. I, 1.8 g. Na2CO3, and 30 cc. H2O refluxed 20 min., the mixture filtered, and acidified to pH 2 with dilute HCl gave 0.7 g. 4-HO2CCH2S analog of XII, needles, m. 221° (decomposition); the filtrate chilled 4 days yielded 0.12 g. [3,4-d]-isomer (XVIII) of XV, m. 305°. XVI (1 g.), 0.9 g. I, 1.8 g. Na2CO3, and 30 cc. H2O refluxed 20 min. and worked up gave 0.45 g. 4-isomer of XVII, m. 173°. 9-Methyl-6-mercaptopurine (1.0 g.) in 10 cc. H2O containing 0.9 g. I and 1.8 g. Na2CO3 refluxed 35 min., the mixture cooled to room temperature, and acidified with dilute HCl gave 1.25 g.

9-methyl-6-carboxymethylthiopurine, m. 225-6° (hot 30% aqueous EtOH). 6-Nitro-4-quinazolone (1.0 g.), 1.5 g. P2S5, and 15 cc. dry C5H5N refluxed 0.5 hr., the whole cooled, poured onto crushed ice, filtered after 2 hrs., and the residue repptd. with AcOH from dilute aqueous NaOH gave 0.93 g. 4-mercapto-6-nitroquinazoline (XIX), bright yellow needles, m. 261-3° (decomposition) (aqueous C5H5N). The 7-NO2 and the 8-NO2 isomers (XX) of XIX, bright yellow needles, m. 270-1° (decomposition) (aqueous C5H5N), and yellow needles, m. 266-7° (decomposition) (aqueous C5H5N), resp., were prepared in 67 and 46%, resp., yields from 7- and 8-nitro-4-quinazolone, resp. 5-Nitro-4-quinazolone (6 g.) and 10.5 g. PC15 heated 3 hrs. at 150°, the mixture cooled, diluted with 150 cc. petr. ether (b. 60-70°), cooled 1 hr. at 0°, and filtered, the residue stirred 10 min. with dilute aqueous NaOH, ice, and CH2Cl2, and the organic layer worked up yielded 4.7 g. 4-chloro-5-nitroguinazoline (XXI), needles, m. 146-7° (sublimed at 130°/0.1 mm.). XXI (1 g.) in 20 cc. dioxane treated with stirring at room temperature with KSH (from 0.3 g. KOH) in 20 cc. absolute EtOH, the whole diluted after 1 hr. with 20 cc. Et.20. and filtered, and the residue added rapidly with stirring to 10 cc. H2O,

and filtered, and the residue added rapidly with stirring to 10 cc. H2O, 0.25 g. NaOH, and 0.4 cc. MeI, and the mixture filtered after 20 min. yielded 0.55 g. 4-methylthio-5-nitroquinazoline, pale yellow flakes, m. 146-7° (petr. ether). XIX (7.35 g.), 400 cc. H2O, 6.8 g. KOH, and 8.4 g. MeI stirred 4 hrs. at room temperature gave 7.2 g. 4-MeS analog (XXII)

of

XIX, m. 162-3°(absolute BtOH). XIX (1 g.), 0.5 g. I, and 20 cc. H2O
refluxed 0.5 hr., the mixture cooled to 0°, and filtered gave 0.43 g.
5-nitroanthranilonitrile (XXIII), m. 210-11° (sublimed at
140°/0.05 mm.). XXII (0.5 g.), 1.24 g. KOH, 40 cc. H2O, and 60 cc.
dioxane stirred 2 hrs. at room temperature, the solution concentrated, and
cooled yielded

0.032 g, XXIII, m, 210°. XX (1 g.) treated with I and K2CO3 in the usual manner gave 0.085 g, 3-isomer of XXIII, yellow needles, m. 137-8° (sublimed at 100°/0.01 mm.).

T 101445-25-4P, Pyrazinamide, 3-amino-5,6-diphenyl-110490-39-6P, Pyrazinamide, 3-amino-5,6-diphenylthio-RL: PREP (Preparation)

(preparation of) RN 101445-25-4 CAPLUS

CN Pyrazinecarboxamide, 3-amino-5,6-diphenyl- (9CI) (CA INDEX NAME)

RN 110490-39-6 CAPLUS

CN Pyrazinamide, 3-amino-5,6-diphenylthio- (6CI) (CA INDEX NAME)

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L7 ANSWER 32 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
                         1958:55949 CAPLUS
DOCUMENT NUMBER:
                         52:55949
ORIGINAL REFERENCE NO.: 52:10106g-i,10107a-i,10108a-i
TITLE:
                        Pteridines. XVI. A synthesis of 2-aminopyrazine-3-
                         carboxamides by reductive ring cleavage of
                         3-hvdroxv-1-pvrazolo[b]pvrazines
AUTHOR(S):
                         Taylor, E. C., Jr.; Barton, J. W.; Osdene, T. S.
                        Princeton Univ., Princeton, NJ
CORPORATE SOURCE:
SOURCE:
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DOCUMENT TYPE:
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OTHER SOURCE(S):
                        CASREACT 52:55949
   cf. C.A. 50, 13047b. PhN:NCH(CN)CO2Et (I) (4.1 g.) and 25 cc. EtOH
     refluxed 15 min. with 1.4 g. N2H4.H2O, cooled to 0°, and filtered
     vielded 3.6 g. 3-hydroxy-4-phenylazo-5-aminopyrazole (II), deep red
     needles, m. 256° (decomposition). HON:C(CN)CONHNH2 N2H4 salt (III) (5.0
     g.) in 25 cc. 40% aqueous NaOH kept 1 hr. at 60°, acidified with
     glacial AcOH, and filtered gave 3.87 g. 3-hydroxy-4-nitroso-5-
     aminopyrazole (IV); a similar run heated 0.5 hr. on the steam bath gave
     2.56 q. IV. III (5.0 q.) in 100 cc. EtOH containing 6 q. Na refluxed 4 hrs.
     with stirring and filtered, and the residue dissolved in 25 cc. H2O,
     acidified with glacial AcOH, and cooled gave 4.0 g. IV. II (4.0 g.) in 50
     cc. 98% HCO2H hydrogenated at 3 atmospheric over 0.4 g. 10% Pd-C, filtered, and
     evaporated, the residue triturated with 1:1 EtOH-Et2O, and the undissolved
     material recrystd. with C from H2O gave 2.95 g. diformyl derivative (V) of
     3-hydroxy-4,5-diaminopyrazole (VI), m. 212-13 (decomposition). IV (2.0
     q.) in 40 cc. 98% HCO2H hydrogenated over 10% Pd-C yielded 2.05 q. V. V
     (8 g.) in 30 cc. 50% H2SO4 warmed to beginning crystallization, diluted with
boiling
     H2O to solution, and cooled slowly yielded 9.4 g. VI.H2SO4, light yellow
     crystals. I (32.5 g.), 7.5 cc. 99% MeNHNH2, and 250 cc. EtOH refluxed 4
     hrs. and cooled to 0° gave 27 g. 1-Me derivative (VII) of II, m.
     265° (EtOH). HON:C(CN)CO2Et (7.1 q.), 5 cc. 99% MeNHNH2, and 30
     cc. EtOH refluxed 3 hrs., refluxed 1 hr. with stirring with 30 cc. 30%
     alc. KOH, cooled to 0°, and filtered, and the residue dissolved in
     20 cc. H2O and adjusted with AcOH to pH 5 vielded 2.9 g. 1-Me derivative
     (VIII) of IV, m. 184-6°; 2nd crop, 0.3 g. VII (20 g.) in 100 cc.
     90% HCO2H hydrogenated 45 min. at 3 atmospheric over 1 g. 10% Pd-C, filtered,
and
     evaporated in vacuo, the residual oil washed with Et20 and dissolved in 70 cc.
     EtOH, and the solution cooled gave 12.8 g. monoformyl derivative (IX) of the
     derivative (X) of VI, m. 210°; it gave recrystd. from aqueous EtOH a
     lower-melting hydrate, m. 188-9° with loss of moisture at
     133-5°. VIII (2.0 g.) in 40 cc. 90% HCO2H hydrogenated in the
     usual manner and evaporated in vacuo, and the residual brown oil dissolved in
     a small amount of EtOH and cooled at 0° yielded 1.5 g. IX, m.
     188-90°. IX (10 g.) recrystd. from 30 cc. 20% H2SO4 containing 25 cc.
     EtOH yielded 13.9 g. X.H2SO4, m. above 300°. 1-Phenyl-3-hydroxy-5-
     aminopyrazole (5.25 g.) in 50 cc. 10% aqueous NaOH added dropwise to PhN2Cl in
     NaOAc buffer (from 3 g. PhNH2, 6 cc. concentrated HCl, 2.1 g. NaNO2, and 12 cc.
     H2O) stirred 0.5 hr., and filtered gave 7.95 g. 1-Ph derivative (XI) of II,
     deep yellow plates, m. 266-8° (decomposition) (Cellosolve).
     2-Phenyl-3-hydroxy-5-aminopyrazole yielded similarly 91% 2-Ph derivative (XII)
     of II, purple-red needles, m. 194-5° (EtOH). I (40 g.), 20 cc.
     PhNHNH2, and 200 cc. iso-AmOH refluxed 24 hrs., cooled to room temperature, and
     filtered, and the residue washed with 100 cc. cold EtOH gave 24.2 g. XII;
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the mother liquor kept at 0° overnight deposited 1.8 g.
phenylazomalonamide phenylhydrazone N-phenylhydrazide, yellow needles, m.
187-8° (EtOH). I (4 g.) and 2 cc. PhNNNNIZ refluxed 20 hrs. with
0.87 g. Na in 75 cc. iso-AmOH and evaporated in vacuo, the residue triturated
with 50% aqueous AcOH, the resulting solid extracted with 200 cc. boiling EtOH,
and the extract concentrated to 50 cc. and cooled yielded 1.39 g. XII; the
EtOH-insol. residue recrystd. from Cellosolve yielded 0.82 g. XI, m.
266-8° (decomposition). XI (5.0 g.) in 50 cc. 90% HCOZH hydrogenated 1
hr. at room temperature and 3 atmospheric over 0.5 g. 10% Pd-C, filtered, and
evaporated in
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on and the oily residue triturated with 50 cc. 1.3 EtOH-Et20 gave 3.1 g. monoformyl derivative (XIII) of 1-phenyl-3-hydroxy-4,5-diaminopyrazole (XIV), plates, m. 223-5° (decomposition) (aqueous EtOH). Crude XIII (3.1 g.) warmed on a water bath with 3 cc. concentrated H2SO4, 7 cc. H2O, and 3 cc. EtOH, diluted with 4 cc. EtOH, and cooled gave 4.8 g. XIV.H2SO4, yellow needles. XII (8.0 g.), 100 cc. 90% HCO2H, and 0.8 g. 10% Pd-C hydrogenated at 3 atmospheric yielded 4.8 g. monoformyl derivative (XV) of 2-phenyl-3-hydroxy-4,5-diaminopyrazole (XVI), m. 235° (decomposition) (aqueous EtOH). XII (12 g.) converted to the XV and the crude product

crystallized

From 1:1 30% H2S04-EtOH yielded 11.6 g. XVI.H2SO4, crange plates.
VI.H2SO4 (20 g.) and 28 g. glyoxal-NaHSO3 adduct (XVII) in 250 cc. H2O treated dropwise with stirring at 60°, stirred 0.5 hr., adjusted to pH 5, cooled to 0°, and filtered gave 9.9 g. 3-hydroxy-1-pyrazolo[b]pyrazine (XVIII), yellow, m. 314-15° (decomposition).
VI.H2SO4 (1.5 g.) in 10 cc. H2O treated with shaking with 1 cc. Ac2 and filtered yielded 0.93 g. 5,6-di-Me derivative (XIX) of XVIII, yellow, m. 325° (decomposition) (sublimed at 230°/0.1 mm.). VI.H2SO4 (4.2 g.), 6.3 g. Bz2, 1.2 g. NaOH, 30 cc. EtOMe, 30 cc. EtOH, and 20 cc. H2O refluxed 1.5 hrs., concentrated in vacuo to about 1/6 its original volume, basified with aqueous NaOH, treated with C, and filtered, the filtrate acidified with HCl, and the precipitate repptd. from aqueous NaOH with HCl and

dried

azeotropically with C6H6 yielded 3.5 g. 5,6-di-Ph derivative (XX) of XVIII, yellow, m. 269° (decomposition) (EtOAc). X.H2SO4 (4.52 g.), 5.6 g. XVII, and 40 cc. H2O adjusted slowly with stirring to pH 5, kept at room temperature overnight, and filtered gave 2.84 g. 1-Me derivative (XXI) of XVIII,

bright yellow needles, m. 242-3° (sublimed at 200°/0.1 mm.).
XVIII (1.0 g.) in 10 cc. 10% aqueous NaOH treated at 60° with stirring
with 1.4 g. MeI and evaporated in vacuo after 45 min., and the residue
dissolved in a little H2O and repptd. With AcOH (pH 5) yielded 0.62 g.
XXI. X.H2SO4 (1.13 g.), 0.5 cc. Ac2, and 10 cc. H2O treated dropwise with
NH4OH to pH 7-8 and readjusted to pH 5 after 10 min. with AcOH gave 0.78
g. 1.5,6-tri-Me derivative of XVIII, m. 268-9° (EtOH and sublimed at
200°/0.1 mm.). X.H2SO4 (1.0 g.), 1 g. Bz2, 10 cc. H2O, 10 cc.
EtAc, and 10 cc. EtOH adjusted to pH 8 with 40% aqueous NaOH, refluxed 1.5
hrs., kept at room temperature overnight, and concentrated in vacuo, the
residue diluted

with H2O, the suspension adjusted with NaOH to pH 9, and the solution heated to boiling, treated with C, filtered, and acidified with AcOH yielded 0.35 g. 1-Me derivative of XX, m. 258-60° (BtOH and sublimed at 200°/0.1 mm.). XVIII (15 g.) in 150 cc. 10% aqueous NaOH and 15 cc. EtCH treated with 15 cc. PhCH2Cl, evaporated after 1 hr. in vacuo, acidified with 50% aqueous AcOH, and filtered gave 18.4 g. 1-PhCH2 derivative (XXII) of XVIII, pale yellow needles, m. 175-6° (MeOH). XIV.H2SO4 (12 g.) and 13 g. XVII in 150 cc. H2O adjusted slowly with concentrated NH4OH to pH

7-8

stirred 45 min., readjusted to pH 5 with glacial AcOH, and cooled to 0° yielded 7.7 g. 1-Ph derivative (XXIII) of XVIII, lime-green needles, m. 227-9° (aqueous EtOH). XVI.H2SO4 (37 g.), 40 g. XVII, and 400 cc. H2O gave in the same manner 23.2 g. 2-phenyl-1-pyrazolo[b]pyrazin-3(2H)-

one (XXIV), pale green plates, m. 232-3.5° (EtOH). XVI.H2SO4 (0.96 g.), 0.4 cc. Ac2, and 100 cc. H2O yielded in the same manner 0.8 g. 5,6-di-Me derivative of XXIV, m. 239-40°, which recrystd. from EtOH and sublimed at 200°/0.1 mm. gave another polymorphic form, m. 193-5°. VI.H2SO4 (8.5 g.) and 8.8 g. NaHSO3 in 100 cc. H2O treated with 6 cc. 47.5% AcCHO, treated dropwise with stirring at 60° until the pH reached 7-8, stirred 45 min., adjusted with dilute AcOH to pH 4-5, and cooled to 0° gave 3.83 g. 6-Me derivative (XXV) of XVIII, light vellow needles, m. 319-21° (H2O); the mother concentrated in vacuo to 1/3 the original volume and kept 24 hrs. at 0° gave 1.15 g. 5-Me derivative (XXVI) of XVIII, buff-colored prisms, m. 234-5° (EtOH). XVIII (1.0 q.), 20 cc. HCONH2, and 3 q. Raney Ni heated 1.5 hrs. with stirring at 115-20°, treated with an addnl. 2 g. catalyst, heated again 1.5 hrs. with stirring, filtered, and cooled yielded 0.58 g. 2-aminopyrazine-3-carboxamide (XXVII), m. 244-5°. XIX (0.5 g.), 50 cc. 95% EtOH, and 6 g. Raney Ni refluxed 2 hrs., filtered, and evaporated, and the solid residue sublimed at 200°/0.1 mm. gave 0.28 g. 5,6-di-Me derivative (XXVIII) of XXVII, light yellow, m. 255°. IV (1.28 g.) in 40 cc. H2O containing 2 cc. concentrated NH4OH refluxed 7 hrs. with 1.2 q. Ac2 and 4 q. Ranev Ni, filtered, and cooled to 0° gave 0.32 q. XXVIII; the Ranev Ni residue extracted with boiling EtOH gave an addnl. 0.06 g. XXVIII. XX (1.0 g.), 50 cc. 95% EtOH, and 8 g. Ranev Ni refluxed 3 hrs., filtered,

and evaporated in vacuo, the residue triturated with H2O and filtered, and the insol. portion washed, dried (0.8 q.), and sublimed at 190°/0.01 mm. yielded the 5,6-di-Ph derivative of XXVII, bright yellow, m. 203-5°. XXI (1.0 g.), 100 cc. 95% EtOH, and 5 g. Ranev Ni refluxed 2.5 hrs., filtered, and evaporated in vacuo gave 0.38 g. 2-MeNH analog of XXVII, light yellow rods, m. 200-1° (sublimed at 180°/0.1 mm.). XXIII (6 g.), 60 g. Raney Ni, and 600 cc. EtOH refluxed 4 hrs. with stirring and filtered through Celite, the filter cake extracted with hot EtOH, the combined filtrate and washing evaporated in vacuo, and the residue (3.2 q.) recrystd. gave the 2-PhNH analog of XXVII, greenish yellow plates from EtOH by slow crystallization or needles by rapid cooling, m. 175-6°. XXIV (5.0 g.), 500 cc. 95% EtOH, and 50 g. Raney Ni refluxed 3 hrs. and filtered, the residue washed with hot EtOH, the combined alc. solns. evaporated, and the residue sublimed at 160-70°/15 mm. yield 52% 2-aminopyrazine-3-carboxylic acid anilide (XXIX), needles, m. 106-7° (EtOH). XXIX (2.0 g.) and 50 cc. 10% aqueous NaOH refluxed 2.5 hrs., diluted with 50 cc. H2O, cooled, and extracted with Et2O, and the aqueous layer adjusted to pH 5 gave 2-aminopyrazine-3-carboxylic acid (XXX), m. 200-1°; the Et20 extract evaporated and the residual oil treated with Ac20 gave 0.41 g. AcNHPh, m. 112-13°. XXII (3.75 g.), 40 g. Raney Ni, and 400 cc. EtOH refluxed 3 hrs. with stirring gave in the usual manner 0.24 g. unchanged XXII and 1.35 g. 2-PhCH2NH analog (XXXI) of XXVII, needles, m. 125-6° (EtOH). XXXI (1.0 q.) and 10 cc. 10% aqueous NaOH refluxed 2 hrs., adjusted to pH 4 with dilute HCl, cooled, and filtered gave 0.78 g. 2-PhCH2NH derivative of XXX, plates, m. 166.5-68° (aqueous EtOH). XXVI (2 g.), 20 g. Raney Ni, and 200 cc. EtOH refluxed 4 hrs. with stirring gave 0.93 g. 5-Me derivative of XXVII, m. 203-4° (MeOH). XXV gave similarly 51.5% 6-Me derivative (XXXII) of XXVII, pale yellow, m. $235-6^\circ$ (sublimed at 160-70°/18 mm.). XXXII (1.0 g.) and 10 cc. 10% aqueous NaOH refluxed 2 hrs., adjusted to pH 4 with dilute HCl, cooled to 0°, and filtered gave 0.72 g. 6-Me derivative of XXX, m. 211-12° (decomposition) (aqueous EtOH).

IT 101445-25-4P, Pyrazinamide, 3-amino-5,6-diphenyl-RL: PREP (Preparation)

⁽preparation of) 101445-25-4 CAPLUS

RN 101445-25-4 CAPLUS CN Pyrazinecarboxamide, 3-amino-5,6-diphenyl- (9CI) (CA INDEX NAME)



m.

L7 ANSWER 33 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1957:76968 CAPLUS

DOCUMENT NUMBER: 51:76968

ORIGINAL REFERENCE NO.: 51:13875a-h
TITLE: Pteridines.

TITLE: Pteridines. V. Derivatives of 1,4-dihydro-1- and 3,4-dihydro-3-methyl-6,7-diphenylpteridine

AUTHOR(S): Boon, W. R.; Bratt, G.

CORPORATE SOURCE: Imp. Chem. Ltd., Manchester, UK
SOURCE: Journal of the Chemical Society (1957) 2159-61

CODEN: JCSOA9; ISSN: 0368-1769

DOCUMENT TYPE: Journal LANGUAGE: Unavailable

AB Condensation of MeHNC(:HH)NH2 with CH2CNCO2Et gave 4-amino-6-hydroxy-2-methylaminopyrimidine and 2,6-diamino-3,4-dihydro-3-methyl-4-oxopyrimidine and not 2,6-diamino-3,4-dihydro-2-methyl-4-oxopyrimidine (Roth, et al., C.A. 46, 3059g). 5,6-Diamino-1,4-dihydro-2-mercapto-1-methyl-4-oxopyrimidine sulfate [1] [Traube and Winter, Arch. Pharm. 244, 16(1906)] (7 g.), 6 g. benzil (II), and 18 g. NaOAc.3H2O (III) refluxed 6 hrs. in 75% aqueous EtOH, the mixture cooled, the product collected, extracted with hot petr. ether (b. 100-20°), and crystallized from BuOH gave 7.4 g. 1,4-dihydro-2-mercapto-1-methyl-4-oxo-6,7-diphenylpteridine (IV), m. 289°. 2,5,6-Triamino-1,4-dihydro-1-methyl-4-oxopyrimidine (6.3 g.), 5.8 g. II, and 17 g. III refluxed 6 hrs. in 25% aqueous EtOH, the solution cooled, the precipitate collected, and crystallized from RCONMe2 (V) gave 10 q.

cooled, the precipitate collected, and crystallized from Hounwez (v) gave 10 g. 2-amino-1,4-dihydro-1-methyl-4- oxo-6,7-diphenylpteridine (VI), m. 333° (decomposition). IV (0.4 g.), 0.5 g. HgO, 70 cc. BuOH, and 10 cc. CHC13 refluxed 6 hrs. in a slow stream of NH3, the mixture filtered hot, the filtrate evaporated in vacuo, and the residue crystallized from V and then from EtOH gave VI, m. 333° (decomposition). 1,4-Dihydro-1-methyl-2-methyl-2-filtered hot, the filtrate from V and then from the control of the composition o

was obtained similarly using MeNH2 in lieu of NH3. VI (0.5 g.) and 50 cc. 2N NaOH refluxed 4 hrs., the solution cooled, acidified with AcOH, the

precipitate
 collected, and crystallized from aqueous EtOH gave 0.16 g.
1,4-dihydro-2-hydroxy-1-

methyl-4-oxo-6,7-diphenylpteridine (VIII), m. 280°. To 0.9 g. I in N KOH was added dropwise with stirring at 100° 01 oc. 1202 (100 volume), the solution cooled, acidified with AcOH, the precipitate (0.3 g.) collected.

and crystallized from EtOH giving VIII, m. 280°. 2-Amino-1,4-dihydro-1-methyl-6,7-diphenyl-4-thionopteridine (IX) (see below) (3 g.) in 300 cc. 2N NaOH refluxed 4 hrs., the solution cooled, acidified, and the product fractionally crystallized from MeOH gave VIII. VI (15 g.), 19.5 g. P2S5, and 300 cc. pyridine (X) refluxed 2 hrs., X removed in vacuo, the residue extracted with 2% aqueous NaOH, and crystallized twice from V gave 7.4 g. IX,

295° (decomposition). On similar treatment, VII gave 16% 1,4-dihydro-1-methyl-2-methylamino-6,7-diphenyl-4-thionopteridine, m. 300° (decomposition) (from V), and IV gave 53% 1,4-dihydro-2-mercapto-1-methyl-6,7-diphenyl-4-thionopteridine, m. 375° (decomposition) (from V without prior extraction with NaOH). 2,4-Diamino-6,7-diphenylpteridine (3 g.), 6 g. MeI, and 60 cc. EKCCHECHECHEOH refluxed 3 hrs., the solution cooled, the

hydriodide [m. 315° (decomposition)] collected, and boiled 5 min, with 10% aqueous Na2CO3 gave 1.7 g. 2-amino-1,4-dihydro-4-imino-1-methyl-6,7diphenylpteridine (XI), m. 256 °. IX (2 g.), 2.5 g. HgO, 120 cc. EtOH, and 20 cc. CHC13 refluxed 6 hrs. in a stream of NH3, the mixture filtered hot, the filtrate cooled, and the product (0.9 g.) crystallized from EtOH gave XI, m. 256°. Similarly was obtained 21% 2-amino-1,4-dihydro-1-methyl-4-methylimino-6,7-diphenylpteridine, m. 256° (from EtOH). 2-Amino-5,6-diphenylpyrazine-3-carboxylic acid (Weijlard, et al., C.A. 39, 30012) Me ester (3.6 g.) and 50 g. MeNH2 in 500 cc. EtOH heated 16 hrs. at 160-70°, the solution cooled, the precipitate collected, and crystallized from MeOH gave 2 g. N:C(NH2).C(CONHMe):NCPh:CPh.N (XII), m. 198°. XII (1.5 g.) and 40 cc. C1CO2Et refluxed 20 hrs., excess C1CO2Et removed in vacuo, and the residue crystallized from CHC13-petr. ether gave 1.7 g. N:C(NHCO2Et).C(CONHMe):N.CPh:CPh.N (XIII), m. 153°. XIII (1.25 g.) refluxed 10 hrs. with NaOMe solution (from 1.5 g. Na in 200 cc. EtOH), EtOH removed in vacuo, the residue suspended in H2O, acidified with AcOH, and the precipitate crystallized from EtOH gave 0.7

g. 3,4-dihydro-2-hydroxy-3-methyl-4-oxo-6,7-diphenylpteridine, m. 307°.

IT 60980-98-5

(Derived from data in the 6th Collective Formula Index (1957-1961))

RN 60980-98-5 CAPLUS

 ${\tt CN \quad Pyrazine carboxamide, \ 3-amino-N-methyl-5,6-diphenyl-\ (9CI) \quad (CA\ INDEX\ NAME)}$

RN

IT 102318-77-4P, Pyrazinecarbamic acid, 3-methylcarbamoyl-5,6-diphenyl-, ethyl ester Ri: PREP (Preparation)

(preparation of) 102318-77-4 CAPLUS

CN Pyrazinecarbamic acid, 3-methylcarbamoyl-5,6-diphenyl-, ethyl ester (6CI) (CA INDEX NAME)

L7 ANSWER 34 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1957:76967 CAPLUS

DOCUMENT NUMBER: 51:76967

ORIGINAL REFERENCE NO.: 51:13870c-i,13871a-i,13872a-i,13873a-i,13874a-i,13875a
TITLE: Pteridines. IV. Derivatives of 2,4-diaminopteridine

and related compounds

Boon, W. R.

AUTHOR(S):

CORPORATE SOURCE: Imp. Chem. Ltd., Manchester, UK
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GI For diagram(s), see printed CA Issue.

AB cf. C.A. 46, 2082g. Several derivs. of 2,4-(H2N)2-Y (in this abstract Y = pteridine) possess antimalarial activity (Potter and Henshall, C.A. 51, 1974h). A series of 2,4,6,7-(H2N)2Ph2-Y were prepared in which the H2N groups were progressively substituted by Me. Antimalarial activity was immediately lost, but the compds. were active against exptl. schistosomiasis in mice. Further modifications of the substituents always lowered the activity. Only a few compds. showed any appreciable activity. 2,4,6-Me2N-(H0)2-Z (in this abstract Z = pyrimidine) ground to pass a 30-mesh sieve, added with stirring during d5 min. to 280 cc. AcOH and 65

schistosomiasis in mice. Further modifications of the substituents always lowered the activity. Only a few compds. showed any appreciable activity. 2,4,6-Me2N-(H0)2-Z (in this abstract Z = pyrimidine) ground to pass a 30-meeh sieve, added with stirring during 45 min. to 280 cc. AcOH and 65 cc. HNO3 (d. 1.5) at 20-5°, stirred an addnl. 45 min., the mixture poured into 1350 cc. H2O, the solid separated, washed free from acid, and dried gave 81 g. 5-O2N derivative (I). I (5 g.), 60 cc. POCl3, and 20 cc. PhNMe2 heated to 105° (bath temperature), after the vigorous reaction the heating continued 1 hr., excess POCl3 removed in vacuo, the residue treated with 200 g. ice, the suspension extracted with four 50-cc. portions of Et20, the combined exts. dried, filtered, evaporated, and the residue

EtZU, the combined exts. dried, filtered, evaporated, an crystallized

from petr. ether (b. 60-80°) gave 3.7 g, 4,6-Cl2 compound (II), m. 117-20°. II (14 g.), 90 cc. C6H6, and 10 cc. aqueous NH3 (d. 0.880) shaken overnight, the mixture filtered, and the residue (4.2 g.) crystallized twice from dioxane gave the 4,6-(H2N)2 compound, m. 249-50°; evaporation of the filtrate gave a residue which, after chromatography on 120 g. Al2O3 in 30 cc. C6H6 and crystallization from EtOAc-petr. ether afforded 0.5 g. 4-H2N compound, m. 132°. To 91 g. Na in 2 l. MeOH was added 509 g. (MeHNC(:NN)NH2]2.H2SO4, the mixture refluxed 30 min. with stirring, CH2(COZE1)2 added, the heating continued 6 hrs., the mixture cooled, diluted with 5 l. H2O. treated with C. filtered, the filtrate acidified to litmus

with 5 1. H2O, treated with C, filtered, the filtrate acidified to litmus with AcOH, and the precipitate collected to give 183 g. 2,4,6-MeHN(HO)2-Z (III); the mother liquors deposited 15 g. presumably 2-amino-1,4,5,6-tetrahydro-1-

the mother liquors deposited 15 g. presumably 2-amino-1,4,5,6-tetrahydro-1-methyl-4,6-dioxo-Z, m. above 360°. III (93g.) and 510 g. POCl3 refluxed 1 hr., the mixture filtered through sintered glass, the filtrate poured on 2250 cc. 32% aqueous NaOH and ice, the separated solid collected, blod

with H2O, and crystallized from MeOH gave 88 g. 2,4,6-(MeIN)(C12-Z (IV), m. 164°. IV (130 g.) heated 12 hrs. with NaOMe (from 168 g. Na in 570 cc. MeOH), the solution cooled, the precipitate collected, washed with H2O, and crystallized from MeOH yielded 95 g. 4,6,2-C1(MeO) (MeN)-Z, m. 153°. Similarly was prepared 81% 4,6,2-C1(MeO) (MeN)-Z (VI), m. 62° (after sublimation at 55°/0.1 mm.), from 4,6,2-C12(MeV2N)-Z at room temperature VI (10 g.) heated 30 min. on a steam bath with 50 cc. HCl, the solution cooled, the product collected, and purified by solution in aqueous alkali, treatment with C, and repptn. with AcOH gave 5.5 g. 6-HO compound, m. 265° (decomposition). Similarly was obtained from VI 95% 4,6,2-C1(H2O) (Me2N)-Z (VII), m. 21°°. 4,6,2-C1(H2O) (Me2N)-Z (VII), m. 21°°. 4,6,2-C1(H2O) (Me2N)-Z (VII) and 78 cc. 19.5% alc. Me2NH heated 17 hrs. at 110-20° gave 172 g. 4-Me2N derivative, m. 172° (from C6H6). Ph(H2N)CHOOPh.HCl (47 g.) dissolved in 750 cc. H2O. basified at 0° with aqueous NH3, the base collected, sucked as dry as possible, added to 35 g. 2,4,6-C13-Z (VIII) in 750 cc. EtOH, the mixture set aside 2 days at room temperature, the precipitate (12 g.)

collected, and crystallized from EtOH gave α -(2,4-dichloro-6-pyrimidylamino)deoxybenzoin (IX), m. 165°. p-ClC6H4CHBZNHZ (X) (28.5 g.) converted to the base, the latter treated as above with 9 g. VIII, the crude product refluxed 3 hrs. with 10 cc. 19.5% alc. Me2NH and

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10 cc. EtOH, the solution evaporated to 0.5 its volume, and the solid recrystd.
     from MeOH gave ω-(4-chloro-2-dimethylamino-6-pyrimidyl-amino)-
     @-(p-chlorophenyl)acetophenone, m. 151-2°; the mother liquors
     gave the 6-Me2N isomer, m. 181-2° (from EtOH), and a small amount of
     another compound believed to be 2,5-di(p-chlorophenyl)-3,6-diphenylpyrazine,
     m. 239-40°. 4,6,2-C12(H2N)-Z (XI) (33 g.) heated 3 hrs. with 175
     cc. 19.5% alc. Me2NH, after the initial reaction had subsided the solution
     cooled, the precipitate (24 q.) collected, and crystallized from MeOH and then
     C6H6 gave 4,2,6-Cl(H2N)(Me2N)-Z, m. 164-5°. Similarly were
     obtained in 70% yield from the appropriate derivative of XI and an alc.
solution
     of H2NCH2CO2Et, Et 4-chloro-2-methylamino-6-pyrimidylaminoacetate (XII),
     m. 167°, and Et 4-chloro-2-dimethylamino-6-pyrimidylamino-acetate,
     m. 121°. 2,4,6-C12(Me2N)-Z (36 g.), 200 cc. EtOH, and 50 cc. 70%
     aqueous EtNH2 refluxed 6 hrs., EtOH removed, the mixture diluted with H2O,
extracted
     with Et20, the extract dried, Et20 removed, the residue dissolved in 70 cc.
     absolute EtOH, 9 cc. concentrated H2SO4 added (the mixture acid to Congo red),
and dry
     Et20 added to a permanent turbidity gave 34 g. 4,6,2-C1(EtNH)(MeNH)-Z
     sulfate, m. 148° (from EtOH-Et20). The following compds. were
     prepared similarly: 4,2,6-Cl(Me2N)(MeNH)-Z, m. 78° (from petr.
     ether); 4.2.6-C1(Et2N)(MeNH)-Z sulfate, m. 148-9° (from Et0H-Et2O);
     4-chloro-6-methylamino-2-piperidino-Z, m. 118° (from MeOH);
     4,6,2-Cl(MeNH)(Me2NCH2CH2NH)-Z, m. 99° (from EtOAc-petr. ether).
     To 17.5 g. VII in 500 cc. H2O containing 60 cc. 2N NaOH and 12.6 g. NaHCO3 was
     added 4-ClC6H4N2Cl (XIII) [from 12.75 g. 4-ClC6H4NH2 (XIV)], the solution
     stirred overnight, the precipitate collected, washed with H2O, EtOH, and Et2O,
     and crystallized from dioxane to give 20 g. 5-p-ClC6H4N2 derivative (XV), m.
     220-2° (decomposition). 4,6,2,5-C1(HO)(MeNH)(p-C1C6H4N2)-Z was obtained
     similarly but could not be purified without decomposition XIII (500 cc.
     0.025M) and 46 g. NaOAc.3H2O (XVI) added with stirring to 3.8 g.
     6,4,2-Me(HO)(Me2N)-Z in 500 cc. H2O, after 16 hrs. the precipitate collected,
     washed, dried in air, and recrystd. from BuOH gave 5.5 g. 5-(p-ClC6H4N2)
     derivative, m. 216-17°. XIII (50 cc. 0.025M) and 40 g. XVI added with
     stirring to 5.0 g. 4,2,6-C1(Me2N)2-Z in 70 cc. AcOH, diluted with 200 cc.
     H2O, after 48 hrs. stirring the solid collected, washed with H2O, and
     crystallized twice from EtOH gave 5 g. 5-(p-ClC6H4N2) derivative (XVII), m.
     91°. The following N.CX:N.CW:C(N:NR).CY (XVIII) (W = C1) were
     prepared (X, Y, R, m.p., crystallization solvent, % vield given): NH2, NHMe,
     p-C1C6H4, 255°, HCONMe (XIX), 47; NH2, NMe2, p-C1C6H4, 204°,
     XIX-EtOH, 65; NHMe, NH2, p-ClC6H4, 272° (decomposition), XIX, 90; NHMe,
     NHMe, p-ClC6H4, 272°, XIX-EtOH, 95; NHEt, NHMe, p-ClC6H4,
     214°, BuOH, 75; NMe2, NH2, p-C1C6H4, 229°, BuOH, 90; NMe2,
     NHMe, Ph, 163°, EtOH, 78; NMe2, NHMe, p-ClC6H4, 183°, BuOH,
     90; HNCH2CH2NMe2, NHMe, p-C1C6H4, 158°, EtOH, 50.
     6,4,2,5-C1(H2N)(Me2N)(p-C1C6 H4N2)-Z (XX) (2 g.) and 40 cc. saturated alc. NH3
     heated 36 hrs. at 150-60°, the solution cooled, and the product (1.75
     q.) crystallized from BuOH gave 6-H2N compound, m. 272-30 [HCl salt, m.
     301° (decomposition) (from 80% HCO2H) (prepared from XIII and
     4,6,2-(H2N)2(Me2N)-Z in AcOH)]. Similarly were prepared the following XVIII
     (W = NH2, R = p-C1C6H4) (X, Y, m.p., crystallization solvent, % yield given):
     NHMe, 213°, BuOH, 40 and 80; NH2, NMe2, 205°, XIX-H2O, 96;
     NH2, NH(CH2)3NEt2, 139°, EtOH-H2O, 44; NHMe, NH2, 241°,
     BuOH, 70; NHMe, NHMe, 197°, EtOAc, 85 and 92; NHMe, NMe2,
     184°, XIX-H2O, 90 and 79; NHEt, NHMe, 161°, BuOH, 80; NMe2,
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NHMe, 193°, BuOH, 90; NMe2, NMe2, 203°, BuOH, 95 and 93; NMe2, piperidino, 175°, BuOH, 86; NMe2, morpholino, 183°, BuOH, 91; NMe2, NH(CH2)2NEt2, 150°, petr. ether, 44; NH(CH2)2NMe2, NHMe, 144°, petr. ether, 90. XVII (5 g.), 100 cc. XIX, and 20 cc.

NH2.

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10% alc. NH3 heated 64 hrs. at 60°, H2O added, and the precipitate crystallized
from EtOH gave 4 g. 4-Me2N derivative (XXI). m. 145°. XXI was also
obtained similarly from XVII and MeOH-Me2NH. Similarly were prepared:
2,4,6,5-(H2N)(Me2N)(MeHN)(p-C1C6H4N2)-Z, m. 192°, and
2,4,6,5-(MeHN)3(p-C1C6H4N2)-Z, m. 155°. 2,4,6,5-(H2N)2(MeHN)(p-C1C
6H4N2)-Z (5 g.) in 75 cc. EtOH reduced by H over Raney Ni (initial
pressure 47 atmospheric) at 90-5° 5 hrs., the mixture acidified with 4 cc.
AcOH, filtered through Hyflo Supercel, the residue washed with H2O, the
combined filtrate and washings evaporated to dryness in vacuo under N, the
residue triturated with Et20, dissolved in 10 cc. H20, acidified to Congo
red with H2SO4, EtOH added, and the precipitate crystallized from H2O gave
2,4,5,6-(H2N)3(MeHN)-Z sulfate (XXII). No satisfactory analytical results
were obtained for 2,5,6,4-(H2N)2(Et2N)(Me2N)-Z oxalate, m. 221°
(decomposition), but it condensed normally with benzil to the pteridine. The
following XC:N.C(NH2):C(NH2).CY:N were prepared (X, Y, m.p., crystallization
solvent, % yield given): NH2, NHMe, 250° (decomposition), H2O, 89; NH2,
NMe2, 209°, aqueous EtOH, 48; NHMe, NH2, 255° (decomposition), H2O,
75; NHMe, NHMe, 259°, aqueous EtOH, 80; NHMe, NMe2, 193°, aqueous
EtOH, 65; NHEt, NHMe, 293° (decomposition), aqueous EtOH, 49; NMe2, NH2, 314° (decomposition), H2O, 58; NMe2, NHMe, 273° (decomposition), H2O,
64; NMe2, NMe2, 182° (decomposition), EtOH, 38; NMe2, piperidino,
208° (decomposition), aqueous EtOH, 33; NMe2, morpholino, 194°
(decomposition), aqueous EtOH, 57. H2NCH2CH(OEt)2 (15 g.) and 17.5 g.
6.4.2.5-Cl(MeHN)-(Me2N)(p-ClC6H4N2)-Z refluxed 24 hrs. in dioxane, the
solution evaporated to dryness, the residue (10 q.) triturated with EtOH,
filtered off, and crystallized from petr. ether gave 5-p-chlorophenylazo-2-
dimethylamino-4-methylamino-6-pyrimidylaminoacetaldehyde di-Et acetal, m.
95°. PhCH(NH2)CH(OMe)2 (XXIII) (11 g.) and XVII in 205 cc. dioxane
refluxed 4 hrs., the solvent removed, and the product (1.9 g.) crystallized
from BuOH gave α-[5-p-chlorophenylazo-2,4-bis(dimethylamino)-6-
pyrimidyl]amino-α-phenylacetaldehyde di-Me acetal, m. 151°.
Similarly was prepared from XV a-(5-p-chlorophenylazo-2-dimethylamino-
4-hydroxy-6-pyrimidyl)-amino-α-phenylacetaldehyde di-Me acetal
(XXIIIa), m. 242° (from BuOH). H2NCH2C(:NNHCONH2)Me.HCl (11 g.)
stirred 2 hrs. with cold NaOEt (from 1.5 g. Na in 60 cc. EtOH), 9.3 g. XV
in 140 cc. XIX added, stirring continued 15 hrs., the semicarbazone, m.
243°, collected, washed with H2O and EtOH, dissolved in 25 cc. AcOH
and 150 cc. 2N aqueous HCl, the solution kept overnight, filtered, the filtrate
evaporated to dryness, and the residue (6.6 g.) crystallized from EtOH gave
5-p-chlorophenylazo-2-dimethylamino-4-hydroxy-6-pyrimidylaminoacetone HCl
salt, m. 217°. The following compds, were prepared similarly:
ω-(5-p-chlorophenylazo-2-dimethylamino-4-hydroxy-6-
pyrimidyl)aminoacetophenone (XXIV) HCl salt monohydrate, m. 229°
(from EtOH) [XXIV semicarbazone, m. 263° (decomposition) (from
XIX-EtOH)]; 4-chloro-ω-(5-p-chlorophenylazo-4-hydroxy-2-methylamino-
6-pyrimidyl)aminoacetophenone (XXIVa), m. 258° (decomposition)
[semicarbazone, m. 264° (from XIX)]; 4'-Cl derivative of XXIV, m.
244° (decomposition) (from XIX-EtOH) [semicarbazone, m. 255°
(decomposition) (from XIX-EtOH)]. IX (17.5 g.) and 60 cc. 2.5M alc. Me2NH refluxed 3 hrs., cooled, the solid (17 g.) collected, dissolved in 200 cc.
AcOH together with 19 g. XVI, a solution of XIII (from 6 g. XIV) added, after
stirring 4 days the resulting precipitate collected, washed with H2O and EtOH,
and crystallized from BuOH gave 10 g. \alpha-(4-chloro-5-p-chlorophenylazo - 2
- dimethylamino-6-pyrimidyl)aminodeoxybenzoin (XXV), m. 254°
(decomposition). XXV (10 g.) refluxed 20 hrs. with 340 cc. 2.5M alc. Me2NH
gave 5.5 g. 4-Me2N derivative, m. 179° (from EtOH). The following
compds. were prepared similarly: ω-(p-chlorophenyl)-ω-(4-chloro-
5-p-chlorophenylazo-2-dimethylamino-6-pyrimidyl)aminoacetophenone, m.
248° (decomposition) (from BuOH), and @-(p-chlorophenyl)-@-
(5-p-chlorophenylazo-2-dimethylamino-6-pyrimidyl)aminoacetophenone, m.
196° (from BuOH). 4-ClC6H4COCH(NH2)Ph.HCl (14.1 g.) dissolved in
800 cc. H2O, made alkaline with aqueous NH3, the base collected, dried over
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added to 7.8 g. XV in 400 cc. XIX, the mixture stirred 24 hrs. at room temperature, the solid collected, and crystallized from XIX-EtoH gave 7 g. 4-chloro-0-(5-p-chlorophenylazo-2-dimethylamino-4-hydroxy-6-pyrimidyl)amino-0-phenylacetophenone, m. 239°. To 5.6 g. HXNCH2CO2Et was added 5.5 g. IX in 150 cc. dioxane, the whole refluxed 8 hrs., cooled, filtered, the filtrate diluted with H2O, the precipitate collected,
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crystallized from BtOAc-petr. ether, and recrystd. from BtOH to give 2 g. Et (4-amino-5-p-chlorophenylazo-2-dimethylamino-6-pyrimidyl)aminoacetate, m. 139°. (For addnl. compds. of this type, cf. Brit. 763,043). Similarly was prepared Et (5-p-chlorophenylazo-2-dimethylamino-4-hydroxy-6-pyrimidyl)-aminoacetate, m. 218°. A solution (1° cc. 0.01 M) of XIII added to 2.5 g. XII in 160 cc. 50% AcOH containing 10 g. XVI, the whole stirred 12 hrs., the precipitate collected, and crystallized from BuOH gave 2

vacuo, the residue triturated with EtOAc, collected, dissolved in H2O, the solution made alkaline with aqueous NH3, and the product (0.1 g.) crystallized from EtOH

gave 2-dimethylamino-7,8-dihydro-4-hydroxy-6-phenyl-Y-0.5 H2O (XXVI), m. 311°, \(2.70 \) m, (Elcm.\)\)\ a 750 in N HCl). Similarly were prepared the following compds: 2,4-bis(dimethylamino)-7,8-dihydro-6,7-diphenyl-Y, m. 278°; 7-p-chlorophenyl-2-dimethylamino-6,7-dihydro-4-methylamino-6-phenyl-Y. m. 267-9° (not analytically pure); 6-p-chlorophenyl-2-dimethylamino-7,8-dihydro-4-hydroxy-7-phenyl-Y HCl salt, m. 346°. XXIVa (2.55 g.) in 300 cc. XIX shaken in H (initial pressure 2 atmospheric) 2 hrs. with 5 g. Raney Ni, the catalyst and XIX removed,

the residue triturated with Et20, the solid collected, and recrystd. from aqueous XIX gave 1.8 g. 6-p-chlorophenyl-2-dimethylamino-7,8-dihydro-4-hydroxy-Y, m. 370°. XXIIIa (5 g.) treated with 10 cc. concentrated HCl in 100 cc. AcOH, after 1 hr. at room temperature H2O added, the precipitate

collected, reduced with H over Raney Ni, the catalyst and solvent removed, the oily residue mixed with 10 cc. AcOH, triturated twice with Et2O, the remaining oil dissolved in 2N HCl, the resulting solid suspended in H2O, treated with dilute aqueous NH3 until the mixture was just alkaline to Brilliant Yellow, the precipitate

(2.3 g.) collected, and crystallized from aqueous XIX gave 7.4.2-Ph(HO)(Me2N)-Y, m.

326° (decomposition), \(\lambda\) 355 m\(\mu\) (ElLm.1\(\text{8}\) 800, in N HCl).

6,4,5,2-HO(H2N)2(Me2N)-Z sulfate (XXVII) (10.7 g.), 6.1 g. PhCOCHO.H2O, 27 g. XVI, and 400 cc. 50\(\text{8}\) aqueous EtOH refluxed 15 min., the mixture cooled, the solid collected, and crystallized from EtOH gave 7.5 g. 6,4,2,5-HO(H2N)(Me2N)(PhCOCHS)-Z, m. 267° (decomposition). Me
3-amino-5,6-diphenylpyrazine-2-carboxylate (1 g.) heated 16 hrs. at
160° with 10 g. MenN2 in 55 cc. EtOH gave 0.5 g.
2-amino-3-N-methylcarbamoyl-5,6-diphenylpyrazine, 197-8° (from
EtOH). 2,4-Disubstituted pteridines were prepared by the following methods
(for addn1. compds., cf. Brit. 763,044, C.A. 51, 13944a): (1) To 0.2 g.
XXVI in 50 cc. 0.5N NaOH was added 0.1 g. KMnO4 in 15 cc. H2O with
stirring over 15 min., after a further 1.5 hrs. EtOH added, MnO2 filtered
off, washed with H2O, the filtrate and washings concentrated to about 50 cc.,
acidified to Congo red with RCl, neutralized with aqueous NH3, and the product

crystallized from EtOH gave 6,4,2-Ph(HO) (Me2N)-Y (XXIX), m. 322° (decomposition), λ 280 (Elcm.1% 910), 355 mm (Elcm.1% 395). (2a) 4,5,2,6-(H2N)2(Me2N)2-Z sulfate (2.94 g.), 6.8 g. XVI, 1.5 g. XXVIII, and 50% aqueous EtOH-refluxed 15 min., the solution cooled, the solid collected, dissolved in 2N AcOH, the solution treated with C, filtered, the filtrate made alkaline with aqueous NH3, and the precipitate crystallized from BuOH and

then from EtOH
gave 7,2,4-Ph(Me2N)2-Y, m. 191°. (2b) XXVII (7.43 g.), 250 cc. 6N
H2SO4, 3.7 g. XXVIII, and 250 cc. EtOH refluxed 2 hrs., EtOH removed in
vacuo, the residual solution cooled in ice, made alkaline with acueous NH3.

filtered,

the filtrate acidified to litmus with dilute $\ensuremath{\mathsf{AcOH}},$ and the precipitate crystallized from

XIX-EtOH gave 6,4,2-Ph(HO)(Me2N)-Y, m. 332°. (2c) XXII (10.8 g.), 14.8 g, benzil, 24 g, XVI, 400 cc. EtOH, and 100 cc. H2O refluxed 5 hrs., the mixture cooled, the precipitate collected, extracted with 0.5N HCl, and the extract

basified with aqueous NH3 gave 6,7,2,4-Ph2(H2N)(Me2N)-Y (XXX), m. 272° (from EtOH). (3) 6,74,2-Ph2(HO)(H2N)-Y (XXXI) (2 g.) and 120 cc. redistd. POC13 refluxed 2 hrs., excess POC13 removed in vacuo, the residue heated 1 hr. with 100 cc. 2.5 M alc. MeNNI2, the alc. removed, the solid extracted with 0.5N HC1, and the extract basified with aqueous NH3 and crystallized from EtOH

gave
 XXX, m. 272°. In a similar series of reactions, XXIX yielded
 6,2,4-Ph(Me2N)2-Y, m. 190°, and 6,4,2-Ph(EtO) (Me2N)-Y, m.
 200° (from EtOH). By using the conditions of Cain, et al. (C.A.
 43, 426ee), there was obtained from XXXI a product (XXXII), m.

6,7,4,2-Ph2(H2N)(Me2N)-

253-9°.

XXXII extracted with 1.5N AcOH left 2-amino-3-N-methylcarbamoyl-5,6-diphenylpyrazine, m. 197-8°; the extract basified with aqueous NH3 and the precipitate crystallized from EtOH gave 6,7,2,4-Ph2 (Me2N) 2-V (XXXIII), m. 266-7°, undepressed with material obtained by condensing 4,5,2,6-(H2N)2 (MeNH) 2-Z with benzil. 6,7,2,4-Ph2 (MS) (HZN)-Y (XXXIV) treated with alc. MeNH3 under the conditions described by Taylor and Cain (C.A. 47, 137h) also gave XXXIII. XXXIV and alc. Me2NH similarly treated gave a product (XXXV), m. 186-215°. XXXV triturated with cold 0.5N AcOH left a residue which, when repeatedly crystallized from MeOH, m. 211°, undepressed with authentic 6,7,2,4-Ph2 (MeNh2-Y obtained by condensing 4,5,2,6-(H2N)2-(Me2N)2-Z with benzil; the acid extract basified with aqueous NH3, and the precipitate crystallized from BoUH gave

Y, m. 236° , undepressed with material obtained by condensing 4,5,6,2-(H2N)3(Me2N)-Z with benzil (4) 7,2,4-Ph(MeHN)2-Y (0.3 g.) and 50 cc. N HCl refluxed 20 hrs., the solution cooled to 50°, made faintly alkaline to Brilliant Yellow with aqueous NH3, the precipitate collected, washed with

H2O, dried, and crystallized from XIX gave 7.4,2-Ph(HO)(MeHN)-Y, m. 387° (decomposition), undepressed with material prepared by 2a, λ 250 mμ (Elcm.1% 700). The following substituted pteridines, N:CX.N:CY.CIC.N.CR.CR.Y:N, were prepared (X, Y, R, R', m.p., crystallization).

olvent,

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214°, EtOH, 2c, 50; NHMe, NHMe, Me, Me, 266°, EtOH, 2c, 28;
     NHMe, NHMe, Ph, H, 264°, XIX, 3, 32; NHMe, NHMe, H, Ph, 256°
     [λ 365 mμ (E1cm.1% 950)], MeOH, 2b, 30; NHMe, NHMe, H,
     p-C1C6H4,294° [λ 365 mμ (E1cm.1% 925)], XIX, 2b, 25;
     NHMe, NHMe, Ph, Ph, 262°, XIX-EtOH, 2c, 49; NHMe, NHMe, o-C1C6H4,
     o-C1C6H4, 265°, BuOH, 2c, 22; NHMe, NHMe, m-C1C6H4, m-C1C6H4,
     256°, MeOH, 2c, 31; NHMe, NHMe, p-C1C6H4, p-C1C6H4, 323°
     XIX, 2c, 63; NHMe, NHMe, p-MeOC6H4, p-MeOC6H4, 259°, EtOH, 2c, 24;
     NHMe, NHMe, 3,4-CH202C6H3, 3,4-CH202C6H3, 297°, XIX-EtOH, 2c, 28;
     NHMe, NHMe, R and R' = 9,10-phenanthrylene, 311°, XIX, 2c, 66;
     NHMe, NHMe, R and R' = 7,8-acenaphthylene, 307°, XIX, 2c, 40; NHMe,
     NHMe, 2-furyl, 2-furyl, 218°, EtOAc, 2c, 24; NHMe, NHMe, R and R' =
     2,3-indolo, 338°, XIX, 2c, 75; NHMe, NMe2, Ph, Ph, 306°,
     XIX, 2c, 60; NHEt, NHMe, Ph, Ph, 249°, EtOH, 2c, 21; NMe2, OH, ph,
     H, 336° (decomposition), EtOH, 1, 2a, and 4, 15 and 90; NMe2, OH, H, Ph,
     325° (decomposition), XIX-EtOH, 1, 2b, and 4, 65, 90, and 90; NMe2, OH,
     p-ClC6H4, H, 377° (decomposition), XIX-EtOH, 1, 85; NMe2, OH, Ph, Ph,
     361°, XIX-EtOH, 2c, 33; NMe2, OH, p-C1C6H4, Ph, 350°, BuOH,
     1, 85; NMe2, OEt, Ph, H, 200°, MeOH, EtOH on 4-Cl compound, 30;
     NMe2, NH2, Ph, Ph, 239°, BuOH, 2c, 63; NMe2, NHMe, Ph, Ph,
     205°, EtOAc, 2c, 43; NMe2, NHMe, Ph, p-C1C6H4, 239° EtOH, 1,
     70; NMe2, NMe2, iso-Pr, iso-Pr, 150°, aqueous EtOH, 2c, 30; NMe2,
     NMe2, Ph. H. 188°, EtOH, 2a and 3, 29 and 40; NMe2, NMe2, H. Ph.
     191°, EtOH, 2b and 3, 37 and 80; NMe2, NMe2, Ph, Ph, 211°
     EtOAc, 2c, 55; NMe2, piperidino, Ph, Ph, 207°, aqueous EtOH, 2c, 75;
     NMe2, morpholino, Ph, Ph, 216°, EtOH, 2c, 71. To a solution of
     PhCH:CHOAc in 290 cc. CC14 was added 39 cc. Br in 40 cc. CC14 with
     stirring below 10° during 1.5 hrs., 290 cc. MeOH added, stirring
     continued 12 hrs. more below 10°, after a further 48 hrs. the mixture
     poured into ice H2O, the separated oil collected, washed with 5% aqueous
NaHCO3,
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dried, and distilled in the presence of a little Na2CO3 to give 122 g. Ph.CHBrCH(OMe)2 (XXXVI), bl4 138-40°. XXXVI (122 g.), 183 g. Ph.CH2NH2, and a trace of NaI heated 1 hr. at 140°, when the reaction had moderated heating continued 2 hrs., the mixture cooled, poured into H2O, the product extracted with Bt2O, the extract dried, and rectified

gave

89 g. PhCH(CH2Ph) CH(OMe)2 (XXXVII), b0.2 121-48°. XXXVII hydrogenated in 300 cc. MeOH over 25 g. 5% Pd-C at 100-5° with an initial pressure of 95 atmospheric, the catalyst removed, and the filtrate rectified gave 47 g. XXIII, b18, 134-6°. BzCH2NH2.HCl (56 g.) dissolved in 350 cc. EtOH with gentle warming, the solution cooled rapidly to room temperature, 25 g. NH2NHCONH2 added, the mixture set aside several hrs.,

the

crystals filtered off, and crystallized from EtOH gave the semicarbazone, m. 107-89. To 28 g. 4-clC6HdCH2Bz in 50 cc. dry Et2O saturated with HCl at 0° was added 7.5 g. BuNO2 in 50 cc. Et2O, the precipitate collected, and crystallized from aqueous MeOH giving the hydroxyimino compound (XXXVIII), m.

121-3°. XXXVIII reduced at room temperature and pressure in 350 cc. EtOH containing 12 cc. concentrated HCl over Pd-C, the catalyst and solvent removed, and

the product (6 g.) crystallized from 2N HCl and then from MeOH-Et2O gave X, m. 248° (decomposition).

60980-98-5P, Pyrazinamide, 3-amino-N-methyl-5,6-diphenyl-

RL: PREP (Preparation) (preparation of)

RN 60980-98-5 CAPLUS

CN Pyrazinecarboxamide, 3-amino-N-methyl-5,6-diphenyl- (9CI) (CA INDEX NAME)

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Ph N C NHMe
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ACCESSION NUMBER:
                           1957:76966 CAPLUS
DOCUMENT NUMBER:
                           51:76966
ORIGINAL REFERENCE NO.:
                          51:13869d-i,13870a-c
TITLE:
                           Syntheses in the quinazolone series. VI. Synthesis of
                           1,2,3,4-tetrahydro-2-aryl-4-oxoquinazolines
AUTHOR(S):
                           Kilroe Smith, T. A.; Stephen, Henry
CORPORATE SOURCE:
                           Univ. Witwatersrand, Johannesburg, S. Afr.
SOURCE:
                           Tetrahedron (1957), 1, 38-44
                           CODEN: TETRAB; ISSN: 0040-4020
DOCUMENT TYPE:
                           Journal
LANGUAGE:
                           Unavailable
OTHER SOURCE(S):
                           CASREACT 51:76966
   cf. C.A. 51, 9626b. N2-Arylideneorthoanilamides (o-
     arylideneaminobenzamides) (I), readily prepared by condensation of aromatic
     aldehydes with o-H2NC6H4CONH2, are characterized by the ease with which
     they isomerize to 1,2,3,4-tetrahydro-2-aryl-4-oxoquinazolines (II). The
     aromatic aldehyde (1 mole) and 1 mole o-H2NC6H4CONH2 refluxed in EtOH, the
     solution cooled, filtered, and the product crystallized from EtOH gave the
     following I (aryl group, m.p., and % yield given): o-HOC6H4, 165°,
     81; o-MeOC6H4, 159°, 77; m-HOC6H4, 146°, 70; p-HOC6H4, 160°, 70; p-MeOC6H4, 158°, 61; 2,4-(HO)2C6H3, 190°,
     90; 2,4-(MeO)2C6H3, 160°, 88; 2,4-(EtO)2C6H3, 177°, 87;
     2,4-EtO(HO)C6H3, 180°, 72; 2,4-HO(EtO)C6H3 (Ia), isomerized, 66;
     3,4-HO(MeO)C6H3 (Ib), 153°, 50; 3,4-MeO(HO)C6H3 (Ic), 187°,
     81; 3,4-EtO(HO)C6H3, 187°, 97; 3,4-(MeO)2C6H3, 165°, 84;
     3,4-EtO(MeO)C6H3, 152°, 60; 2,3-HO(MeO)C6H3, 168°, 81;
     o-O2NC6H4, 174°, 86; m-O2NC6H4, 199°, 95; p-O2NC6H4,
     191°, 93; PhCH:CH, 210°, 90; and 2,3,4-HO2C(MeO)2C6H2,
     208°, 96. Ia, Ib, and Ic isomerized during recrystn, from EtOH and
     were alkylated for identification and analysis. The I refluxed 30 min.
     with N HCl, then with 2N NaOH containing EtOH, or heated above the m.p. in
     vacuo in some instances gave good yields of the II [aryl, m.p., and %
     yield from the acid (a), base (b), or by heating (c) given]: Ph,
     228°, -; p-MeC6H4, 230°, -; o-HOC6H4, 300°, 82a;
     m-HOC6H4, 209°, 100b; p-HOC6H4, 332°, 70a; o-MeOC6H4,
     181°, 88b; p-MeOC6H4, 195°, 62a; 2,4-HO(EtO)C6H3,
     305°, 100c; 2,4-(EtO)2C6H3, 149°, 94b; 2,4-(MeO)2C6H3,
     187°, 100b; 2,3-HO(MeO)C6H3, 279°, 87a; 3,4-MeO(HO)C6H3, 224°, 92a; 3,4-HO(MeO)C6H3, 191°, -; 3,4-EtO(MeO)C6H3,
     89°, -, 3,4-Eto(HO)C6H3, 218°, -, 3,4-(MeO)2C6H3, 226°, 100b; o-O2NC6H4, 192°, 96b; PhCH:CH, 294°, 58b; 3,4-(CH2O2)C6H3, 202°, -; 2,3,4-HO2C(MeO)2C6H2, 296°, 100b,
     100c. II in dry Me2CO treated in a period of 2-3 hrs. with KMnO4 in dry
     Me2CO, the excess KMnO4 removed with NaHSO3, filtered, the Me2CO evaporated,
     and the residue crystallized from MeOH or EtOH gave 2-aryl-4-quinazolinones
     (III) (aryl, m.p., and % yield given); Ph (IIIa), 238°, 70;
     p-MeC6H4 (IIIb), 241°, 73; p-MeOC6H4, 208°, 50; p-MeOC6H4,
     247°, 98; o-O2NC6H4, 237°, 95; m-O2NC6H4, 354°, 96;
     p-02NC6H4, 365°, 90; 2,4-(MeO)2C6H3, 204°, 75;
     2,4-(EtO)2C6H3, 174°, 87; 3,4-(MeO)2C6H3, 247°, 65;
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ANSWER 35 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

3,4-(CH2O2)C6H3, 279°, 75; 3,4-EtO(MeO)C6H3, 239°, 90; PhCH:CH, 252°, 44 (cf. Stephen and Wadge, C.A. 51, 6649e). BzH (10.6 g.) and 15.1 g. o-H2NC6H4CO2Me in petr. ether (b. 60-80°) kept 3 days at 0° (CO2 atmospheric) and the product (75%) crystallized from petr. ether (b. 40-60°) gave o-PhCH(OH)NHC6H4CO2Me (IV), m. 77°. Similar condensation with p-MeC6H4CHO gave the corresponding o-[4-MeC6H4CH(OH)NH]C6H4CO2Me (IVa), m. 79°. IV and IVa kept 2 weeks at 0° in EtOH saturated with NH3 gave 41% IIIa and 58% IIIb. BzH (4 g.) and 10 g. o-H2NC6H4CO2Me warmed in 50 cc. EtOH containing a trace of HCl, and the orange solution refluxed 40 min, and filtered hot gave 8.6 g. white solid, m. 265-75°, yielding on extraction with Me2CO 6.9 g. insol. 1,2,3,4-tetrahydro-3-(o-carbomethoxyphenyl)-4-oxo-2-phenylquinazoline and 1.7 g. Me2CO-soluble (o-MeO2CC6H4NH)2CHPh, m. 188-90°. Refluxing 10.3 g. o-H2NC6H4CO2H and 12.5 g. 2,4-H0(EtO)C6H3CHO in EtOH gave 19.8 g. 2-[o-2,4-HO(EtO)C6H3CH:N]C6H4CO2H, m. 206°. Similarly were prepared the corresponding 2,4-EtO(HO) and 2,3-HO(MeO) analogs, m. 211° and 119°, in 97 and 80% yields, resp. 60980-98-5 (Derived from data in the 6th Collective Formula Index (1957-1961))

Pyrazinecarboxamide, 3-amino-N-methyl-5,6-diphenyl- (9CI) (CA INDEX NAME)

0 || Ph N C-NHMe

60980-98-5 CAPLUS

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AB

L7 ANSWER 36 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1957:9378 CAPLUS

ACCESSION NUMBER: 1957:9378 CAPLUS DOCUMENT NUMBER: 51:9378

ORIGINAL REFERENCE NO.: 51:1971b-e

TITLE: A new synthetic approach to pteridines

AUTHOR(S): Osdene, T. S.; Taylor, E. C.
CORPORATE SOURCE: Princeton Univ., Princeton, NJ

SOURCE: Journal of the American Chemical Society (1956), 78,

1-methyl-3-hydroxy-1-pyrazolo[b]pyrazine (IX), m. 242-3°. The

5451-2

CODEN: JACSAT; ISSN: 0002-7863
DOCUMENT TYPE: Journal

LANGUAGE: Journal Unavailable

cf. C.A. 50, 13047b. A general method is described for the synthesis of pyrazine intermediates which permits the ready synthesis of 1-substituted pteridines. PhN2CH(CN)CO2Et with N2H4 or N2H4.H2O in EtOH yielded 3-hydroxy-4-phenylazo-5-aminopyrazole (I), m. 256° (decomposition). I with H in 98 HCO2H containing 10% Pd-C yielded 3-hydroxy-4,5-5 diformylaminopyrazole (II), m. 212-13° (decomposition). II with 50% H2SO4 yielded 3-hydroxy-4,5-5-diaminopyrazole sulfate (III). Cyclization of the N2H4 salt of nitrosocyanoacetohydrazide with 40% NaOH at room temperature yielded 3-hydroxy-4-nitroso-5-aminopyrazole (IV); catalytic reduction of IV yieldsd III. The same reactions with MeNENH2 yielded 1-methyl-3-hydroxy-4,5-diaminopyrazole, m. above 250°. III with glyoxal, Ac2, and Bz2 yielded 3-hydroxy-1-pyrazolo (b) pyrazine (V), m. 314-15° (decomposition); 3-hydroxy-5,6-dimethyl-1-pyrazolo)[b]pyrazine (VII), m. 325° (decomposition); 3-hydroxy-5,6-dipenyl-1-pyrazolo)[b]pyrazine (VII), m. 269° (decomposition); 1-methyl-3-hydroxy-5,6-dimethyl-1-pyrazolo)[b]pyrazine (VIII), m.

preceding compds. treated with Raney Ni yielded 2-amino-3-carboxamides. VII treated with Raney Ni 3 hrs. in boiling EtOH yielded 80% 2-amino-5,6-diphenylpyrazine-3-carboxamide, m. 203-5°. Similarly, IX vielded 2-methylaminopyrazine-3-carboxamide, m. 200-1°. Direct condensation of IV with Ac2 in EtOH containing Ranev Ni vielded 2-amino-5,6-dimethylpyrazine-3-carboxamide. 101445-25-4P, Pyrazinamide, 3-amino-5,6-diphenyl-

RL: PREP (Preparation)

(preparation of)

RN 101445-25-4 CAPLUS CN

Pyrazinecarboxamide, 3-amino-5,6-diphenyl- (9CI) (CA INDEX NAME)

ANSWER 37 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1956:69468 CAPLUS

DOCUMENT NUMBER: 50:69468

ORIGINAL REFERENCE NO.: 50:13047b-i,13048a-b

TITLE: Pteridines. XIV. Further studies on a new approach to

pteridine synthesis

AUTHOR(S): Taylor, E. C., Jr.; Garland, Robert B.; Howell,

Charles F.

CORPORATE SOURCE: Univ. of Illinois, Urbana

SOURCE: Journal of the American Chemical Society (1956), 78, 210-13

CODEN: JACSAT; ISSN: 0002-7863 DOCUMENT TYPE: Journal

LANGUAGE: Unavailable OTHER SOURCE(S): CASREACT 50:69468

cf. C.A. 50, 2608h. 3-Amino-5,6-diphenylpyrazinamide (I) (1.509 g.) and 10 cc. BzCl refluxed 4 h., cooled, and diluted with 250 cc. petr. ether gave

1.179 g. 2,6,7-triphenvl-4(3H)-pteridinone (II), white needles, m. 290° (from CH2Cl2-petr, ether and then aqueous HCONMe2) (all m.ps. are corrected). The N-PhCH2 derivative (III) of I (0.5 g.) and 25 cc. AcCl

refluxed 4

h. and diluted with 25 cc. petr. ether yielded 0.36 g. 3-acetylamino-5,6diphenylpyrazinamide (IV), bright vellow platelets, m. 207-8° (from CHCl3-petr. ether). III (0.835 g.), 10 cc. Ac20, and 10 cc. MeCN refluxed 4 h. and evaporated to dryness in vacuo, and the residue treated with EtOH and evaporated to dryness again gave 0.472 g. N-PhCH2 derivative (V) of IV, tan crystals, m. 149-50° (from CH2Cl2-petr. ether). V (0.613 q.) refluxed 3 h. with 0.5 g. Na in 10 cc. absolute EtOH and poured into 50 cc. H2O gave 0.503 g. III, m. $186-7^{\circ}$. 3-PhCH2 derivative of II gave similarly 93% III. I (2.53 g.), 5 cc. PhNCO, and 25 cc. dry pyridine refluxed 1 h. and cooled yielded 2.81 g. 3-(3-phenylureido)-5,6diphenylpyrazinamide (VI), light yellow platelets, m. 240.5-1.5°

(from aqueous EtOH and then aqueous HCONMe2). III (0.80 g.), 1 cc. PhNCO, and 10

cc. dry pyridine refluxed 2 h., cooled, treated with C, and diluted with petr. ether gave 1.03 g. N-PhCH2 derivative (VII) of VI, sparkling white platelets, m. 210° (from aqueous EtOH). VI (0.523 g.) and 7 g.

polyphosphoric acid (VIII) heated 2 h. at 150° (CO2 was evolved), and diluted with 50 cc. H2O, and the precipitate sublimed at 200° and 2 mm.

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gave 0.134 g. I, m. 204-5°; the sublimation residue sublimed at
300° and 2 mm. gave 3,5,7-tripheny1-2,4(1H,3H)-pteridinedione (IX),
colorless solid, m. 327-8° (decomposition). III and VIII heated 45 min.
at 150° gave 52% I and 63% VII. I (0.97 g.), 2 cc. PhNCO, and 10
cc. pyridine refluxed 3 days, cooled, diluted with 40 cc. CH2Cl2 and 250 cc.
petr. ether, and filtered, and the filtrate evaporated to dryness gave 0.418
q. IX, white needles, m. 327-8° (decomposition) (from aqueous HCONMe2). III
gave similarly 51% IX. I (1.52 g.), 3 cc. PhNCS, and 15 cc. pyridine
refluxed 1 h., cooled, and diluted with 150 cc. petr. ether yielded 1.92 q.
3-(3-phenylthioureido) analog (X) of I, light yellow platelets, m.
233° (from aqueous HCONMe2). I (1.67 g.), 3 cc. PhNCS, and 15 cc.
pyridine refluxed 3 days, cooled overnight, and filtered gave 1.87 g.
2-mercapto-3,6,7-triphenyl-4(3H)-pteridinone (XI), fine yellow needles, m.
301-2° (sublimed at 250° and 1 mm.). X heated similarly
with PhNCS gave also XI. N-Bu derivative of I (2.70 g.), 3.5 cc. PhNCS, and
10 cc. pyridine refluxed 4 days, cooled, and diluted with 20 cc. CH2C12 and
100 cc. petr. ether yielded 1.49 g. 2-PhNH analog of XI, pale yellow
crystals, m. 323-4° (from aqueous HCONMe2). I (1.34 g.), 2 cc.
iso-PrNCS, and 20 cc. pyridine refluxed 2 days, cooled, and diluted with 20
cc. CHCl3 and 100 cc. petr. ether gave 1.05 g. 3-(3-isopropylthioureido)
analog (XII) of VI, white platelets, m. 251-2° (from
CH2Cl2-cyclohexane). III (1.04 g.), 1.2 cc. iso-PrNCS, and 15 cc.
pyridine refluxed 2 days and poured onto 200 g. ice vielded 0.7 g. N-PhCH2
derivative (XIII) of XII, pale yellow crystals, m. 170° (from 70%
AcOH). XII (1.24 q.) refluxed 6 h. with 1 q. Na in 25 cc. absolute EtOH,
poured into 100 cc. H2O, and filtered, and the orange solid digested with
dilute HCl gave 0.174 g. 2-mercapto-3-isopropyl-6,7-diphenyl-4(3H)-
pteridinone, light yellow needles, m. 270° (from aqueous EtOH); the
filtrate acidified with concentrated HCl gave 0.72 g. 2-isopropylamino-6,7-
diphenyl-4(3H)-pteridinone (XIV), bright lemon-yellow platelets, m.
324-5° (from aqueous EtOH). XIII (0.390 g.) refluxed 3 h. with 0.1 g.
Na in 5 cc. absolute EtOH and poured into 50 cc. H2O yielded 0.30 g. 3-PhCH2
derivative of XIV, sparkling yellow crystals, m. 305-7° (decomposition)
(from aqueous HCONMe2). 3-Amino-5,6-diphenylthiopyrazinamide (XV) (1.1 g.)
and 10 cc. BzCl refluxed 1.5 h., cooled, diluted with 50 cc. EtOH, refluxed
1 h., and evaporated to dryness, and the residue suspended in hot EtOH and
filtered gave 2,6,7-triphenyl-4(3H)-pteridinethione, yellow crystals, m.
323-4° (sublimed). XV (1.23 g.), 3.4 cc. PhNCS, and 10 cc.
pyridine refluxed 2 h., cooled, and diluted with 180 cc. petr. ether yielded
2.06 q. compound C47H33N9O (structure tentatively assigned), fine yellow
needles, m. 369-70° (from aqueous HCONMe2), also obtained by refluxing
the mixture for 3 days. It was recovered in 93% yield after refluxing 43 h.
with concentrated HCl. XV (1.04 g.), 2 cc. PhNCS, and 10 cc. pyridine refluxed
36 h., diluted with 150 cc. hot petr. ether, and allowed to stand gave a
small amount of unidentified, colorless needles, m. 72-157°, fine
yellow needles, and cushions of orange prisms. The fine yellow needles and orange prisms recrystd. from pyridine-petr. ether yielded 1.15 g.
2-anilino-6,7-diphenvl-4(3H)pteridinethione, long vellow needles, m.
261-2°.
7596-73-8P, Pyrazinamide, 3-amino-N-benzyl-5,6-diphenyl-
857180-32-6P, Urea, 1-[3-(benzylcarbamoyl)-5,6-diphenylpyrazinyl]-
3-phenyl- 857180-53-1P, Pyrazinamide, 3-acetamido-N-benzyl-5,6-
diphenyl- 857183-71-2P, Urea, 1-(3-carbamoyl-5,6-
diphenylpyrazinyl)-3-phenyl-2-thio- 857993-08-9P, Urea,
1-[3-(benzylcarbamoyl)-5,6-diphenylpyrazinyl]-3-isopropyl-2-thio-
859297-19-1P, Pyrazinamide, 3-acetamido-5,6-diphenyl-
859300-58-6P, Pyrazinamide, 3-(3-isopropyl-2-thioureido)-5,6-
diphenyl- 859300-59-7P, Urea, 1-(3-carbamoyl-5,6-
diphenylpyraziny1)-3-phenyl-
RL: PREP (Preparation)
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CN Pyrazinecarboxamide, 3-amino-5,6-diphenyl-N-(phenylmethyl)- (9CI) (CA INDEX NAME)

- RN 857180-32-6 CAPLUS
- CN Pyrazinamide, N-benzyl-5,6-diphenyl-3-(3-phenylureido)- (5CI) (CA INDEX NAME)

- RN 857180-53-1 CAPLUS
- CN Pyrazinamide, 3-acetamido-N-benzyl-5,6-diphenyl- (5CI) (CA INDEX NAME)

- RN 857183-71-2 CAPLUS
- CN Pyrazinamide, 5,6-diphenyl-3-(3-phenyl-2-thioureido)- (5CI) (CA INDEX NAME)

CN Pyrazinamide, N-benzyl-3-(3-isopropyl-2-thioureido)-5,6-diphenyl- (5CI) (CA INDEX NAME)

- RN 859297-19-1 CAPLUS
- CN Pyrazinamide, 3-acetamido-5,6-diphenyl- (5CI) (CA INDEX NAME)

- RN 859300-58-6 CAPLUS
- CN Pyrazinamide, 3-(3-isopropyl-2-thioureido)-5,6-diphenyl- (5CI) (CA INDEX NAME)

- RN 859300-59-7 CAPLUS
- CN Pyrazinamide, 5,6-diphenyl-3-(3-phenylureido)- (5CI) (CA INDEX NAME)

L7 ANSWER 38 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1956:52652 CAPLUS DOCUMENT NUMBER: 50:52652

50:10103e-a ORIGINAL REFERENCE NO.:

TITLE:

Route to 4-aminopteridines

AUTHOR(S):

Taylor, E. C., Jr.; Paudler, W. W.

CORPORATE SOURCE: Princeton Univ., Princeton, NJ

Chemistry & Industry (London, United Kingdom) (1955) SOURCE:

1061-2

CODEN: CHINAG; ISSN: 0009-3068

DOCUMENT TYPE: Journal Unavailable

LANGUAGE:

OTHER SOURCE(S): CASREACT 50:52652

A new route for 4-amino-5,6-diphenylpteridines (I) is described.

2-Hydroxy-5,6-diphenylpyrazinamide (II) (Jones, C.A. 43, 3009h) gave 99% yield 2-chloro-3-cyano-5,6-diphenylpyrazine (III) when heated in a sealed tube with PC13. III was also obtained in 80% yield by heating a mixture of II, POC13, and PC15. Fusion of III with guanidine carbonate, urea, or thiourea gave 65, 59, and 51% 2-amino, 2-hydroxy, and 2-mercapto derivs. of I, resp. III with N2H4.H2O gave 2-chloro-5,6-diphenylpyrazinoic acid hydrazide, or when repeated in the presence of KI gave 3-amino-5,6-diphenyl-1-pyrazolo[b]pyrazine. III gave 2-amino-5,6diphenylpyrazinamide when treated with NH4OH and KI, or 2-amino-3-cyano-5,6-diphenylpyrazine when fused with NH4OAc.

101445-25-4P, Pyrazinamide, 3-amino-5,6-diphenyl-

RL: PREP (Preparation)

(preparation of) RN

101445-25-4 CAPLUS Pyrazinecarboxamide, 3-amino-5,6-diphenyl- (9CI) (CA INDEX NAME) CN

AUTHOR(S):

L7 ANSWER 39 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1954:25073 CAPLUS DOCUMENT NUMBER: 48:25073

ORIGINAL REFERENCE NO.: 48:4553h-i,4554a-i,4555a-d

TITLE: Pteridines, X. A new approach to the synthesis of pteridines

Taylor, E. C., Jr.; Carbon, John A.; Hoff, Dale R.

CORPORATE SOURCE: Univ. of Illinois, Urbana Journal of the American Chemical Society (1953), 75,

1904-8

CODEN: JACSAT: ISSN: 0002-7863

DOCUMENT TYPE: Journal

LANGUAGE: Unavailable OTHER SOURCE(S): CASREACT 48:25073

For diagram(s), see printed CA Issue.

cf. C.A. 48, 2719c. A new synthesis of pteridines is described involving the preliminary synthesis of a 2,4(1H,3H)-pteridinedione (lumazine) by the conventional method and the subsequent aminolytic cleavage of the pyrimidine portion of the lumazine to give a 3-amino-N-substituted

pyrazinamide, followed by its ring closure to the desired pteridine. This method permits a much wider variation in the structure of the pyrimidine ring than does the conventional approach. Dry freshly distilled BuNH2 (100 cc.) and 15 g. 6,7-diphenyl-2,4(1H,3H)-pteridinedione (I) heated 12 h. in a sealed tube at 180°, the clear light brown solution treated with Norit, the excess BuNH2 removed in vacuo, and the residue diluted with 50 cc. hot EtOH and then hot H2O to incipient crystallization gave 8.8 g. (53.3%) 3-amino-N-butyl-5,6-diphenylpyrazinamide (II), bright yellow prisms, m. 146-7° (from CHC13-aqueous EtOH). 3-Amino-N-benzvl-5,6diphenylpyrazinamide (0.520 g.) in 20 cc. HC(OEt)3 (III) and 20 cc. Ac20 refluxed 5 h., and the solution evaporated to dryness in vacuo yielded 0.386 g. (72.3%) 3-benzyl-6,7-diphenyl-4(3H)-pteridinone (IV), white platelets, m. 248° (from CHCl3-petr. ether). II (1.0 g.) in 20 cc. 98-100% HCO2H and 20 cc. Ac20 refluxed 5 h., and the clear light yellow solution evaporated repeatedly to dryness in vacuo with 50-cc. portions of EtOH gave 0.337 g. (32.8%) 3-Bu analog (V) of IV, white platelets, m. 194-5° (from CHCl3-aqueous EtOH). II (0.50 g.), 20 cc. III, and 20 cc. Ac20 refluxed 5 h. similarly gave 0.396 g. (77%) V. 3-Amino-N-benzyl-5,6diphenylpyrazinamide (1.0 g.) and 25 cc. ClCO2Et (VI) refluxed 20 h., and the resulting clear yellow solution evaporated repeatedly to dryness with

50-cc.

portions of EtOH gave 0.996 g. (93.7%) N-benzvl-3-carbethoxyamino-5,6diphenylpyrazinamide (VII), colorless prisms, m. 129-30° (from CHC13-petr. ether). II (2.0 g.), and 40 cc. VI refluxed 20 h. gave similarly 1.539 g. (63.7%) N-Bu analog (VIII) of VII, colorless prisms, m. 110-11° (from CHCl3-petr. ether). VII (0.574 g.) and alc. NaOEt (from 0.5 g. Na in 70 cc. absolute EtOH) refluxed 20 h. gave 0.211 g. (40.9%) 3-benzyl-6,7-diphenyl-2,4(1H,3H)pteridinedione (IX), long colorless needles, m. 194-5° (from CHCl3-petr. ether). VIII (1 g.) similarly gave 0.80 g. (88.8%) 3-Bu analog of IX, long white needles, m. 246-7° (from CHCl3-petr. ether). 3-Amino-N-benzyl-5,6diphenylpyrazinamide (X) (0.597 g.) and 25 cc. HCONH2 heated 3 h. at 190°, and the mixture cooled and diluted with H2O yielded 0.304 q. (64%) 6,7-diphenyl-4(3H)-pteridinone (XI), m. 297-8° (from aqueous HCONMe2), also obtained by refluxing X with HCONH2 containing 2 cc. dilute HCO2H. II similarly gave 52% XI. Me 3-amino-5,6-diphenylpyrazinoate (0.856 g.) in 75 cc. MeOH saturated with anhydrous NH3 at 0° and heated 1 h. at 120° in a sealed tube yielded 0.700 g. (86%) 3-amino-5,6-diphenylpyrazinamide (XII), m. 204-5° (from aqueous EtOH). XII (0.529 g.), 1.0 g. P2S5, and 15 cc. dry pyridine refluxed 1 h., the deep red solution cooled, poured into 200 cc. H2O, the resulting orange colloidal suspension dissolved by the addition of a small amount of 10% NaOH, the solution treated with C, filtered, and the filtrate acidified with glacial AcOH gave 0.304 g. (54.6%) 3-amino-5,6-diphenylthiopyrazinamide (XIII), orange needles, m. 158-60° (from aqueous EtOH). XI (2.975 q.), 4 g. P2S5, and 50 cc. anhydrous pyridine refluxed 2 h. similarly gave 2.34 g. (75%) 6,7-diphenyl-4(3H)-pteridinethione (XIV), bright red platelets, m. 270-80° (decomposition) (from aqueous HCONMe2). XIII (0.286 g.) in 10 cc. III and 10 cc. Ac20 refluxed 5 h. gave 0.164 g. (55.4%) XIV, bright red shiny platelets. XIV (0.5 g.), 1 cc. PhCH2NH2, 1 g. HgO, and 30 cc. EtOH refluxed 5 h., the mixture filtered, the black residue washed with 10 cc. hot EtOH, and the filtrate combined with the washings and diluted with H2O until crystallization began vielded 0.61 g. (99%) 4-benzylamino-6,7diphenylpteridine (XV), light yellow platelets, m. 178-9° (from aqueous Me2CO). XIV (0.951 g.), 1.5 cc. BuNH2, 1 g. HgO, and 20 cc. absolute EtOH refluxed 2.5 h. similarly gave 0.870 g. (74.3%) N-Bu analog (XVI) of XV, bright yellow plates, m. 150-1° (from aqueous EtOH). XIV (2.0 g.) and 50 cc. absolute EtOH saturated with NH3 at 0° and heated in a sealed tube 10 h. at 130° gave 1.59 g. (84%) 4-amino-6,7-diphenylpteridine, light yellow needles, m. 175° (from aqueous Me2CO). Refluxing 0.924 g. XIV in 5 cc. CHC13 and 20 cc. absolute EtOH with 0.8 g. HgO yielded 0.414 g. (33%) mercuric salt of XIV, light yellow crystals, m. 268-71° (from

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CHCl3-absolute EtOH). XV (0.20 g.) in 10 cc. 6N HCl refluxed 0.5 h. and the
cooled mixture neutralized with NH4OH gave 0.14 g. (93%) XI, m.
297-8°. XI (88%) was also formed by hydrolysis of XVI. II (1.75
g.), 2.0 g. P2S5, and 25 cc. dry pyridine refluxed 1 h., the mixture cooled,
poured into 150 cc. H2O, and the precipitate washed with H2O and recrystd. from
absolute EtOH gave 1.54 q. (83.4%) 3-amino-N-buty1-5,6-
diphenylthiopyrazinamide (XVII), bright yellow needles, m. 168-9°.
XVII (0.635 g.), 0.7 g. freshly fused NaOAc, 10 cc. 98-100% HCO2H, and 10
cc. Ac20 refluxed 5 h. gave 0.441 g. (67.6%) 3-butv1-6,7-diphenv1-4(3H)-
pteridinethione (XVIII), orange needles, m. 193-5° (from
CHC13-EtOH). XVII (1.53 g.) in 10 cc. HC(OEt)3 and 10 cc. Ac20 refluxed 3
h. yielded 0.962 g. (61.28) XVIII. XVII (1.139 g.) in 30 cc. ClCO2Et
refluxed 20 h., the solution evaporated to dryness in vacuo, and the residue
evaporated 3 times with 50-cc. portions of absolute EtOH yielded 1.11 g. (77%)
carbethoxy derivative (XIX), microcryst. orange solid, m. 173-4° (from
CHCl3-EtOH). XIX heated 15 min. with 5 cc. 10% aqueous NaOH in 20 cc. EtOH
gave 73% 1,2-dihydro-2-oxo derivative of XVIII, orange-red solid, m.
205-9° (from aqueous EtOH). XVIII (0.179 q.) in 1.5 cc. CHC13 and 10
cc. absolute EtOH refluxed 6 h. with 0.2 g. HgO while a continuous stream of
NH3 was passed through the mixture, the mixture filtered hot, and the filtrate
evaporated to a small volume deposited 0.119 g. (69.8%) 3-butyl-4(3H)imino-6,7-
diphenylpteridine, yellow platelets, m. 149-51°.
3-Amino-5,6-diphenylpyrazinoic acid piperidide (1.50 g.) in 50 cc. VI
refluxed 5 h. and the mixture worked up in the usual manner gave 1.42 g.
(79%) 3-carbethoxyamino-5,6-diphenylpyrazinoic acid piperidine (XX),
yellow platelets, m. 174-5° (from aqueous Me2CO and then CH2Cl2-petr.
ether). XX (0.50 g.) in 40 cc. EtOH saturated with dry NH3 and heated 6 h. in
a sealed tube at 155°, the solution evaporated to dryness, the residue
dissolved in dilute NH4OH, and the solution acidified with glacial AcOH gave
0.330 g. (90%) I, colorless microcryst. solid, m. 320-5°.
7509-57-1P, Pyrazinamide, 3-amino-N-butyl-5,6-diphenyl-
101445-25-4P, Pyrazinamide, 3-amino-5,6-diphenyl-
110490-39-6P, Pyrazinamide, 3-amino-5,6-diphenylthio-
857180-46-2P, Pyrazinamide, 3-amino-N-butyl-5,6-diphenylthio-
857992-95-1P, Pyrazinecarbamic acid, 3-(benzylcarbamoyl)-5,6-
diphenyl-, ethyl ester 857993-29-4P, Pyrazinecarbamic acid,
3-(butylcarbamoyl)-5,6-diphenyl-, ethyl ester
RL: PREP (Preparation)
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7509-57-1 CAPLUS CN

RN

(preparation of)

Pvrazinecarboxamide, 3-amino-N-butvl-5,6-diphenvl- (9CI) (CA INDEX NAME)

101445-25-4 CAPLUS RN

CN Pyrazinecarboxamide, 3-amino-5,6-diphenyl- (9CI) (CA INDEX NAME)

RN 110490-39-6 CAPLUS

CN Pyrazinamide, 3-amino-5,6-diphenylthio- (6CI) (CA INDEX NAME)

RN 857180-46-2 CAPLUS

CN Pyrazinamide, 3-amino-N-butyl-5,6-diphenylthio- (5CI) (CA INDEX NAME)

RN 857992-95-1 CAPLUS

CN Pyrazinecarbamic acid, 3-(benzylcarbamoyl)-5,6-diphenyl-, ethyl ester (5CI) (CA INDEX NAME)

RN 857993-29-4 CAPLUS

L7 ANSWER 40 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 1954:14779 CAPLUS

DOCUMENT NUMBER: 48:14779

ORIGINAL REFERENCE NO.: 48:2719b-e

TITLE: Pteridines. IX. Hydrolytic ring cleavage of 3-benzyl-6,7-diphenyl-4(3H)-pteridinone

AUTHOR(S): Taylor, E. C., Jr.

CORPORATE SOURCE: Univ. of Illinois, Urbana
SOURCE: Journal of the American Chemical Society (1952), 74,

2380-1 CODEN: JACSAT: ISSN: 0002-7863

DOCUMENT TYPE: Journal

LANGUAGE: Unavailable

cf. C.A. 48, 688c, 689g. 5,6-Diamino-4-hydroxy-2-mercaptopyrimidine (15.0 g.) in 300 cc. boiling water dissolved by the addition of 20% Na2CO3, the pH adjusted to 10 with dilute HCl, 80 g. wet Raney Ni added portionwise, the mixture refluxed 4 hrs., cooled, filtered, treated with 12.4 g. Bz2 in 100 cc. MeCOEt and 350 cc. EtOH, refluxed 8 hrs., acidified, and cooled vielded 13.2 g. 6,7-diphenvl-4(3H)-pteridinone (I), m. 297-8° (decomposition). I (0.5 g.), 30 cc. MeOH, 0.2 cc. PhCH2Cl, and 0.16 g. KOH refluxed 2 hrs., and the mixture treated with 15 cc. 2 N NaOH and warmed yielded 0.483 g. 3-amino-N-benzyl-5,6-diphenyl-4-pyrazinamide (II), m. 188.5-89°. 3-Benzyl-6,7-diphenyl-4(3H)-pteridinone (III) in 30 cc. MeOH treated 0.1 g. KOH in 5 cc. water, and the mixture refluxed 10 min. and diluted with 5 cc. water yielded 64 mg. II, m. 188.5-89°. I (1.0 g.), 0.186 g. KOH, 3.8 cc. PhCH2Cl, and 30 cc. MeOH refluxed 1 hr., and the mixture treated with 3 cc. AcOH and hot water to incipient crystallization yielded 0.26 g. III, m. 248°; dilution of the EtOH filtrate yielded 0.19 g. II, m. 187°; the mother liquor on dilution with 1 volume water yielded 0.195 g. I. In another experiment refluxing 24 hrs. yielded 0.21 g. III, m. 248°. 7596-73-8P, Pyrazinamide, 3-amino-N-benzyl-5,6-diphenyl-

1 /596-/3-8F, Fyrazinamide, 3-amino-N-Denzyi-5,6-dipnenyi-RL: PREP (Preparation)

(preparation of) RN 7596-73-8 CAPLUS

CN Pyrazinecarboxamide, 3-amino-5,6-diphenyl-N-(phenylmethyl)- (9CI) (CA INDEX NAME)

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L7 ANSWER 41 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN
ACCESSION NUMBER:
                         1954:3618 CAPLUS
DOCUMENT NUMBER:
                         48:3618
ORIGINAL REFERENCE NO.: 48:688c-i.689a
                         Aminolysis of heterocyclic amides. I. The aminolysis
TITLE:
                         of 6,7-diphenyllumazine
AUTHOR(S):
                         Taylor, E. C., Jr.
CORPORATE SOURCE:
                         Univ. of Illinois, Urbana
SOURCE:
                         Journal of the American Chemical Society (1952), 74,
                         1651-5
                         CODEN: JACSAT; ISSN: 0002-7863
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         Unavailable
    cf. following abstract An alkylamine with 6,7-diphenyllumazine (I) gives
     first an N-substituted amide of a 3-(3-alkylureido)-5,6-diphenylpyrazinoic
     acid, which can then be converted to an N-substituted amide of
     3-amino-5,6-diphenylpyrazinoic acid by further reaction with the amine.
     The mechanism of these transformations is discussed and the results are
     interpreted as a substantiation for the ring cleavages previously
     postulated (cf. C.A. 47, 137h) in the reaction of 4-NH2 and
     4-hydroxy-2-mercaptopteridines with alkylamines. I (3.0 g.) in 20 cc.
     PhCH2NH2 (II) refluxed 15 min. and diluted with 50 cc. absolute EtOH vielded
     2.18 g. N-benzyl-3-(3-benzylureido)-5,6-diphenylpyrazinamide (III). EtOH,
     m. 88-93°; III m. 150-1°. III (0.60 g.), 10 cc. Ac20, and 3
     q. NaOAc refluxed 2 h., and the cooled mixture poured on ice and let stand
    overnight yielded III, m. 150-1°. III (0.50 g.) in 10 cc. II refluxed 8 h., diluted with 20 cc. EtOH, heated to boiling and diluted with
     water to incipient precipitation yielded 0.348 g. 3-amino-N-benzyl-5,6-
     diphenylpyrazinamide (IV), m. 188.5-9°; the filtrates from IV
     concentrated to 20 cc. and diluted with 20 cc. water yielded N,N'-dibenzylurea
     (V), 168°. I and II refluxed 8 h. yielded directly IV and V.
     H2SO4 (2 cc.) added slowly to 1.0 g. 3-amino-5,6-diphenylpyrazinoic acid
    in 15 cc. absolute EtOH, the solution let stand 24 h. at room temperature, and
poured
     into 75 cc. water yielded 0.91 g. Me ester (VI), m. 204-6°. VI
     (165 mg.) and 2 cc. II refluxed 10 min., diluted with 15 cc. 50% EtOH and
     cooled yielded 190 mg. IV, m. 188.5-89°. IV (1.0 g.), 20 cc. 85%
     HCO2H, 20 cc. Ac2O, and 1.0 g. NaOAc refluxed 5 h. and the solution evaporated
† o
     dryness vielded 0.42 3-benzyl-6,7-pteridin-4(3H)-one, m. 248°. I
     (0.50 g.) and 15 cc. morpholine refluxed 14 h. yielded 0.53 g.
     3-(morpholinocarbonylamino)-5,6-diphenylpyrazinoic acid morpholide (VII),
    m. 262-4°. VII (1.0 q.) sealed in 20 cc. morpholine heated 12 h.
     at 140° and 6 h. at 190° yielded 0.64 g.
     3-amino-5,6-diphenylpyrazinoic acid morpholide (VIII), m. 190.5-1°.
     I and morpholine heated 12 h. at 190° yielded VIII directly. I
     (3.0 g.), 30 cc. piperidine, and 10 cc. HCONMe2 refluxed 16 h., filtered,
     and the hot filtrate treated with boiling water to incipient turbidity
     yielded 1.67 g. 3-(piperidinocarbonylamino)-5,6-diphenylpyrazinoic acid
     piperidide, m. 215-17°. I (5.0 g.) in 50 cc. piperidine heated 20
     h. at 200° yielded 3.8 g. 3-amino-5,6-diphenylpyrazinoic acid
    piperidide, m. 156°. I (0.50 g.) in 15 cc. HOCH2CH2NH2 refluxed 12 h. yielded 0.453 g. 3-amino-N-(\beta-hydroxyethyl)-5,6-
     diphenylpyrazinamide, m. 186.5-87°. I (2.0 g.) and 40 cc. NH40H
     heated 16 h. at 185° yielded 1.67 g. 3-amino-5,6-
     diphenylpyrazinamide (IX), m. 203.5-5°. IX (0.3 g.) and 1 cc. II
     refluxed 15 min., diluted with 10 cc. EtOH, and hot water added to incipient
     crystallization yielded 0.31 g. IV. IX (0.06 g.), 5 cc. piperidine, and 2 cc.
     HCONMe2 refluxed 16 h. yielded 0.053 g. IX, m. 203.5-5°.
     p-02NC6H4NHCONH2 (2.0 g.) and 20 cc. piperidine refluxed 8 h. yielded 2.43
     g. 1-(p-nitrophenyl)-3-(piperidino)urea, m. 165-6°. I (1.0 g.) and
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10 cc. 85% H4N2.H2O refluxed 6 h. and the mixture let stand 3 h. at 0° yielded 0.705 g. 3-amino-5,6-diphenylpyrazinoic acid hydrazide

(X), m. 250-1°. The mother liquors from X evaporated to dryness, the residue washed with water, dried, extracted with CH2C12, and the extract diluted

with petr. ether yielded 3-amino-6,7-diphenyl-2,4(1H,3H)-pteridinedione, m. 259-60° (decomposition); evaporation of the filtrates yielded about 0.015

- IT 7509-58-2P, Urea, 1-benzyl-3-[3-(benzylcarbamoyl)-5,6-diphenylpyrazinyl]- 7596-73-8P, Pyrazinamide, 3-amino-N-benzyl-5,6-diphenyl- 101445-25-4P, Pyrazinamide, 3-amino-5,6-diphenyl- 857180-39-3P, Ethyl alcohol, compound with
 - N-benzyl-3-(3-benzylureido)-5,6-diphenylpyrazinamide 857183-65-4P, Pyrazinamide, 3-(2-hydroxyethylamino)-5,6-diphenyl-857984-47-5P, Pyrazinoic acid, 3-amino-5,6-diphenyl-, hydrazide
 - RL: PREP (Preparation)
 - (preparation of)
- RN 7509-58-2 CAPLUS
- CN Pyrazinecarboxamide, 5,6-diphenyl-N-(phenylmethyl)-3-
 - [[[(phenylmethyl)amino]carbonyl]amino]- (9CI) (CA INDEX NAME)

- RN 7596-73-8 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5,6-diphenyl-N-(phenylmethyl)- (9CI) (CA INDEX NAME)

- RN 101445-25-4 CAPLUS
- CN Pyrazinecarboxamide, 3-amino-5,6-diphenyl- (9CI) (CA INDEX NAME)

- RN 857180-39-3 CAPLUS
- CN Pyrazinamide, N-benzyl-3-(3-benzylureido)-5,6-diphenyl-, compd. with EtOH (5CI) (CA INDEX NAME)

CM 1

CRN 7509-58-2 CMF C32 H27 N5 O2

Ph N C-NH-CH₂-Ph O NH-C-NH-CH₂-Ph

CM 2

CRN 64-17-5 CMF C2 H6 O

н₃C[−] Сн₂[−] Он

RN 857183-65-4 CAPLUS

CN Pyrazinamide, 3-(2-hydroxyethylamino)-5,6-diphenyl- (5CI) (CA INDEX NAME)

RN 857984-47-5 CAPLUS

CN Pyrazinoic acid, 3-amino-5,6-diphenyl-, hydrazide (5CI) (CA INDEX NAME)

$$\begin{array}{c|c} \text{Ph} & & \\ \text{Ph} & & \\ \text{N} & & \\ \text{N} & & \\ \text{NH}_2 & & \\ \text{O} & & \\ \end{array}$$

L7 ANSWER 42 OF 42 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1949:15234 CAPLUS

DOCUMENT NUMBER: 43:15234

ORIGINAL REFERENCE NO.: 43:3009e-i,3010a

TITLE: Pyrazines and related compounds. I. A new synthesis of hydroxypyrazines

AUTHOR(S): Jones, Reuben G.

SOURCE: Journal of the American Chemical Society (1949), 71,

78-81 Journal

CODEN: JACSAT; ISSN: 0002-7863 A general synthesis of 2-hydroxypyrazines (I) involves the condensation of

DOCUMENT TYPE: LANGUAGE:

Unavailable

1,2-di-CO compds. with a-amino acid amides. H2NCH2CONH2 and (CHO)2 give 48% I, m. 187-9°. dl-Methionine Et ester (II) (287 g.) in 2 l. absolute EtOH, saturated at 0° with NH3 and kept 30 days, gives 175 g. (93% on basis of unrecovered II) dl-methioninamide (III), m. 48-9°. α-Amino-α-phenylacetamide (IV), m. 128-9°. H2NCH(CONH2)2 (V) (117 g.), added to 25 g. 40% aqueous (CHO)2 diluted with 25 mL. H2O, the mixture treated (temperature below 10°) with 10 mL. 12.5 N NaOH and, after several hrs., with 10 mL. AcOH, give 90% of the 3-carbamyl derivative of I, m. 265° (decomposition); a higher temperature or less (CHO)2 gives a smaller yield; KOH or Et2NH can be used in place of NaOH. AcCHO (36 g.) in 50 mL. H2O at -20°, treated with 60 g. V and then (dropwise, temperature below 0°) with 40 mL. 12.5 N NaOH, kept 18 h. at room temperature, and acidified with 50 mL. 12 N HCl, gives 59% 2-hydroxy-3-carbamyl-5-methylpyrazine (VI), m. 243-4° (decomposition); Ac2 gives 93% of the 5,6-di-Me analog (VII), m. 231-2° (decomposition). V (11.7 g.) and 21 g. Bz2 in 350 mL. 50% aqueous EtOH at 70°, treated with 10 mL, 12.5 N NaOH, give 83% of 2-hydroxy-3-carbamy1-5,6diphenylpyrazine, m. 174-5°; 5-Ph analog m. 213-16°, 75%. 3-Me derivative of I m. 140-2°, 83.7%; 3,5-di-Me derivative m. 145-6°, 42% from MeCH(NH2)CONH2 and AcCHO; 3-methyl-5-Ph derivative m. 212-13°, 56.5%; 5,6-di-Ph derivative m. 225-7°, 97%; 5,6-di-Me derivative m. 199-200°, 11.3%. II and Ac2 in CHCl3 containing 1 equivalent piperidine give 70% (NaOH gives 88%) of the 3-(2-methylmercaptoethyl)-5,6dimethyl derivative of I m. 128-9°; 3-(2-methylmercaptoethyl) derivative of I m. 96-7°, 97%. 3-Ph derivative of I m. 172-3°, 88.5%;

3-phenyl-5,6-dimethyl derivative of I m. 222-6°, 45%. p-HOC6H4CH2CH(NH2)CONH2 and (CHO)2 give 76% of the 3-(p-hydroxybenzyl) derivative of I, m. 212-13°; AcCHO gives 47% of the

3-(p-hydroxybenzyl)-5-Me derivative, m. 202-3°; Ac2 gives 77.5% of the 3-p-hydroxybenzyl-5,6-dimethyl derivative, m. 236-7°. VII (11.5 g.) in 75 mL. 3 N NaOH, heated several hrs. on the steam bath, gives 79% 2-hydroxy-5,6-dimethyl-3-pyrazinoic acid, m. 172-4° (decomposition); VI gives 30% of the 5-Me analog, m. 155-7° (decomposition); the 6-Me isomer, tan, m. 183-4° (decomposition).

34121-79-4P, Pyrazinamide, 3-hydroxy-5,6-diphenyl-RL: PREP (Preparation)

(preparation of)

34121-79-4 CAPLUS

CN Pyrazinecarboxamide, 3,4-dihydro-3-oxo-5,6-diphenyl- (9CI) (CA INDEX NAME)

RN

=> log hold\ 'HOLD\' IS NOT VALID HERE For an explanation, enter "HELP LOGOFF".
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 SINCE FILE
 TOTAL

 COST IN U.S. DOLLARS
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 FULL ESTIMATED COST
 222.28
 420.67

 DISCOUNT AMOUNTS (FOR QUALIFYING ACCOUNTS)
 SINCE FILE
 TOTAL

 CA SUBSCRIBER PRICE
 -32.76
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